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Hanley, ME, Bouma, TJ and Mossman, HL ORCID logoORCID:
<https://orcid.org/0000-0001-5958-5320> (2019) The gathering storm: optimiz-
ing management of coastal ecosystems in the face of a climate-driven threat.
Annals of Botany, 125 (2). pp. 197-212. ISSN 0305-7364

Downloaded from: <https://e-space.mmu.ac.uk/625211/>

Version: Published Version

Publisher: Oxford University Press (OUP)

DOI: <https://doi.org/10.1093/aob/mcz204>

Please cite the published version

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The gathering storm: optimizing management of coastal ecosystems in the face of a climate-driven threat

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The gathering storm: optimizing management of coastal ecosystems in the face of a climate-driven threat

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Running Head: Coastal plants and extreme storm events

ABSTRACT

- **Background** The combination of rising sea levels and increased likelihood of extreme storm events poses a major flood and erosion threat to our coastlines. As a result, many ecosystems recognized and valued for their important contribution to coastal defence, face increased damage from erosion and flooding. Nevertheless, only recently have we begun to examine how plant species and communities, respond to, and recover from, the many disturbances associated with storm events.

- **Scope** We review how the threats posed by a combination of sea level rise and storms affects coastal sub-, inter-, and supra-tidal plant communities. We consider ecophysiological impacts at the level of the individual plant, but also how ecological interactions at community-level, and responses at landscape-scale, inform our understanding of how and why an increasing frequency and intensity of storm damage is vital to effective coastal management. While noting how research is centred on the impact of hurricanes in the US Gulf region, we take a global perspective and consider how ecosystems worldwide (e.g., seagrass, kelp forests, sand dunes, saltmarsh, mangroves) respond to storm damage and contribute to coastal defence.

- **Conclusions** The threats posed by storms to coastal plant communities are undoubtedly severe, but beyond this obvious conclusion, we highlight four research priority areas. These call for studies focusing on (1) how storm disturbance affects plant reproduction and recruitment; (2) plant response to the multiple-stressors associated with ACC and storm events; (3) the role of ecosystem-level interactions in dictating post-disturbance recovery; and (4) models and long-term monitoring to better predict where and how

storms and other climate change-driven phenomenon impact coastal ecosystems and services. In so doing, we argue how plant scientists must work with geomorphologists and environmental agencies to protect the unique biodiversity and pivotal contribution to coastal defence delivered by plant communities.

Key Words: Coastal Erosion - Flooding – Hurricanes -Kelp – Mangrove – Pine savannah - Salt Marsh – Sand Dunes – Seagrass - Sea-Level Rise - Storm Surge – Wave Attenuation

INTRODUCTION

The past, present, and likely future impacts of Anthropogenic Climate Change (ACC) on terrestrial plant species and communities are widely reported and reasonably well understood (Parmesan and Hanley, 2015). Most studies focus on long-term, chronic effects, but considerable environmental threat is likely to stem from an increased frequency and intensity of acute, extreme events (Vasseur *et al.*, 2014; Parmesan and Hanley, 2015). Although chronic stressors doubtless reduce ecosystem resilience, for many coastal plant communities the most important manifestation of ACC is likely to come from the acute disturbance, erosion, and flooding associated with storm events.

In their most recent assessment of our changing climate, the Intergovernmental Panel on Climate Change (IPCC 2019) asserted that anthropogenically-driven Sea Level Rise (SLR), in tandem with an increase in storm frequency and intensity, poses a severe environmental threat to estuarine and coastal ecosystems (ECEs). Nonetheless, plant biologists have recognized this threat only recently, and when combined with our inability to predict where and when storms might occur, it is perhaps no surprise that relatively few authors have systematically addressed the issue. In-fact much of the initial relevant research was conducted in the SE United States where low-lying freshwater wetlands regularly experience periodic seawater inundation as a result of isostatic movements and subsidence, and changes in channel flow regime. Studies by Haller *et al.* (1974), McKee and Mendelssohn (1989) and Flynn *et al.* (1995) reporting species-specific variation in Floridian and Louisianan freshwater marsh plants to ‘natural’ salinity pulses, were

78 nonetheless prescient of how these communities can be expected to respond to
79 contemporary and predicted changes in frequency and intensity of ACC-linked extreme
80 events. Subsequently, a body of work conducted around the Gulf of Mexico has described
81 the responses of wetland vegetation to the disturbance associated with recent hurricanes
82 (Tate and Battaglia, 2013; Meixler, 2017; Imbert, 2018).

83 The realization that coastlines globally now face increasing erosion and flood risk provides
84 the impetus for understanding how hurricanes, typhoons, cyclones and other extreme
85 weather events affect coastal vegetation. Moreover, in many vulnerable locations, ECEs
86 have ‘added value’ in that they offer natural coastal protection against erosion and flooding
87 (Temmerman *et al.*, 2013; Morris *et al.*, 2018). This key ecosystem service has
88 considerable socio-economic benefits, reducing flood risk and damage for a fraction of the
89 costs associated with constructing so-called ‘hard defences’ like concrete walls (Narayan *et al.*,
90 2016; Morris *et al.*, 2018). Nonetheless, society is only just beginning to appreciate this
91 valuable service and how ECEs can be integrated into a dynamic flood defence strategy.
92 Consequently, understanding the response of vegetation to shifts in storm regimes is critical
93 to ensure effective risk management over coming decades.

94 With this mind, we offer here a synthesis of the response of ECE vegetation to extreme
95 storm events, and signpost how an understanding of these responses aids management of
96 ECEs for flood and erosion mitigation. We contextualize recent scientific studies by
97 exploring the threats to, and response of, plants challenged by both SLR and increasing
98 storm frequency and severity. This necessitates understanding ecophysiological responses
99 from the level of the individual, up to geomorphological factors operating across the entire

tidal range. We highlight also future research priorities, from laboratory experiments to large-scale modelling and mapping of post-disturbance vegetation responses, needed to provide an appreciation of the wider ecosystem services delivered by coastal habitats. By bringing together this diversity of topics, our aim is not only to signpost interdisciplinary research towards better management of ECEs, but also promote their integration into strategic coastal defence.

THREATS TO COASTAL ECOSYSTEMS

Although historically, land use change, pollution, and invasive species have all impacted ECEs, and while these threats are certain to continue into the future, our focus is on ACC. Indeed, there seems little doubt that ACC will pose the greatest challenge to coastal habitats for the remainder of this century and beyond (Millennium Ecosystem Assessment, 2005).

Although elevated atmospheric CO₂ (eCO₂), and associated shifts in temperature, and precipitation will have profound effects on all plant communities (Parmesan and Hanley, 2015), the combination of SLR, and increased sea surface temperatures (SST) and enhanced wave forcing is a particular pressing and unique issue for ECEs.

Rising sea levels have already affected many coastal regions. IPCC (2019) stated with ‘*high confidence*’ that the 0.32m increase in global sea levels observed between 1970-2015 was attributable to ACC-driven thermal expansion of the seas and glacier mass loss. It seems clear that SLR will accelerate into the 21st century, although IPCC (2019) have ‘*high confidence*’ that variation in ocean dynamics and coastal land-use will generate regional departures of about 30% around global averages. Not only does this place coastal regions

and habitats at significant (but varying) flood risk, there is ‘*high confidence*’ that SLR will continue for centuries, even if global mean temperatures are stabilized (IPCC, 2019). The ramifications of these changes are severe. IPCC (2019) has ‘*very high confidence*’ that low-lying coastal areas will increasingly experience submergence, flooding and erosion throughout this century and beyond.

It is important however, to distinguish between the impacts of long-term, chronic changes in Earth’s climate, and those imposed by acute ACC-linked events. Although an annual maximum predicted global SLR of 15 mm yr⁻¹ (IPCC 2019) poses problems for coastal plants due to landward/upward displacement of the freshwater-saltwater aquifer interface (White and Kaplan, 2017), SLR and extreme weather together are likely to deal the greatest environmental threat to our coastlines (IPCC, 2019). A combination of increased SST coupled with SLR, is widely predicted to increase the frequency, severity and geographical distribution of tropical cyclones and storm surge events (IPCC, 2019). Consequently, present-day ‘one per century’ sea level extremes are expected on an annual basis for most coastlines by 2100 (IPCC, 2019). Not only will many supra-tidal ECEs face an increased risk of short-duration, seawater inundation as a result, the wave energies and sediment disturbance associated with intense storm activity will impact the many ECEs that help protect coastlines. In addition, most coastal habitats are strongly inter-connected, such that acute erosion and sediment loss from one (e.g. a sub-tidal sand bar), has major repercussions for sediment transport to nearby supra-tidal habitat (e.g. sand dunes) (Hanley *et al.*, 2014).

Indeed, where sufficient ‘pre-event’ data are available, studies show major changes in coastal geomorphology and vegetation for many years afterwards. Carter *et al.* (2018) for example, used a time series of remotely sensed images to show major breaching, land-area reduction, and vegetation loss throughout the Mississippi-Alabama barrier islands in the first 10 months after Hurricane Katrina made landfall. These changes were however, site-specific depending on sediment removal or accretion, underscoring the more general problem that it is difficult to predict exactly how and when storms affect particular coastlines. For example, in the unusually energetic series of winter storms that affected SW England in 2013/4, the most severe impacts coincided with high spring tides and occurred on west-facing beaches where subsequent dune erosion was extensive (Masselink *et al.*, 2015). Similarly, variation in wind directions meant a brackish marshland in Louisiana, USA, apparently unaffected by Hurricane Katrina in August 2005, experienced major seawater incursion following Hurricane Rita only a month later (Steyer *et al.*, 2007).

The spatio-temporal stochasticity associated with forecasting storm events presents a major limitation to our ability to predict where and when ECEs will be impacted. Nevertheless, it seems certain that ECEs globally can expect a significant increase in erosion and flood frequency and duration over coming decades. In Table 1, we summarize how the threats associated with extreme storms are likely to affect coastal habitats across the tidal range, and in the following sections, discuss how some of these key threats, exert major ecological effects on sublittoral, inter-tidal, and supra littoral habitats.

IMPACTS ON COASTAL PLANT COMMUNITIES

Supra-tidal Plant Communities

Vegetation subject to seawater immersion at exceptionally high tides or during storm surge events only. Affected habitats include sand dunes, and other (semi-)natural terrestrial and aquatic ecosystems (grasslands, pine savannah, freshwater wetlands).

Due, in part, to our inability to predict where and when storm surges will occur, and even less effectively, control and replicate natural flood events, few field studies deal with the impact of storm disturbance on supra-tidal plant communities. Although remote sensing offers a way to assess and monitor largescale changes in vegetation following storm events (e.g. Carter *et al.*, 2018; Douglas *et al.*, 2018; Stagg *et al.*, 2020), elucidating how saltwater flooding, mechanical damage, litter accumulation, and sediments affect the plant community is challenging. There is however, a relatively large body of research describing the (species-specific) effects of burial by sediments on sand dune species (Sykes and Wilson, 1990; Harris *et al.*, 2017; Brown and Zinnert, 2018), while Tate and Battaglia, (2013) and Platt *et al.*, (2015) report major negative effects of simulated post-hurricane litter deposition on Floridian and Mississippian pine savannah. Surprisingly however, few studies consider the immediate effects of physical damage on supra-littoral coastal vegetation (see Platt *et al.*, 2000).

The most widely reported impact of ACC-linked extreme events on supra-littoral ECEs is seawater flooding. Immersion in seawater brings additional problems for supra-littoral plants compared to those experienced by species in inland riparian, or coastal inter-tidal

communities. Flooding of the former is exclusively freshwater, while plants in most inter-tidal ECEs have an inherent ability to tolerate salinity associated with (twice-daily) tidal immersion. Although by virtue of their association with the coast, sand dune, cliff edge, and other supra-littoral plants may be tolerant of salt spray (Malloch *et al.*, 1985; Sykes and Wilson 1988), the combination of anoxia and salt stress imposed by seawater flooding is unique to these habitats.

In fact the ‘salt stress’ associated with coastal flooding seems to be much more important to plant response and recovery than anoxia. In experiments where supra-littoral plants have been simultaneously exposed to freshwater and seawater immersion, the former has never resulted in any noticeable impact on plant ecophysiology compared with untreated (no immersion) controls (Tolliver *et al.*, 2009; Hanley *et al.*, 2013, 2017, 2020a,b; White *et al.*, 2014). A full appraisal of how and why salinity stress affects plant ecophysiology is beyond the scope of this review (see instead Flowers and Colmer, 2008; Munns and Tester, 2008; Negrão *et al.*, 2017; the latter an excellent assessment of methods to evaluate plant physiological responses to salinity stress). In short however, high seawater salinity (of which chloride (55%) and sodium (31%) contribute most of the ‘salt’ content), causes both osmotic (limiting the plant's ability to absorb water) and ionic (increased toxicity via Na⁺ and Cl⁻ accumulation) stresses (Munns and Tester, 2008). It is worth bearing in mind though that our oceans have marked seasonal and regional salinity variation (Donguy and Meyers, 1996) and that seawater is much more than ‘NaCl in solution’. Some ions such as K⁺ and Ca²⁺ have direct negative toxicological or osmotic effects, but also the potential to mitigate the impact of Na⁺ and Cl⁻ on plant metabolism (Flowers and Colmer, 2008; Munns

and Tester, 2008). It is likely that other ions have similar moderating influences over Na⁺ and Cl⁻ stress, and consequently, understanding how seawater affects plant ecophysiological responses requires much more than a simplistic evaluation of the effects of NaCl alone. This point was reinforced by **Hanley *et al.*, (2020a)**, who show how short-duration immersion of *Trifolium repens* in NaCl solutions elicited almost total mortality compared to plants subject to immersion in natural seawater or commercially available marine aquarium salt solutions.

It is possible to monitor ECE recovery after a natural flood event (e.g. Flynn *et al.*, 1995; Lantz *et al.*, 2015), but this requires the ability to allocate resources quickly to an affected site in order to capture changes in vegetation as floodwaters recede. Moreover, to appreciate fully post-inundation transitions, a thorough understanding of the pre-flood ecosystem is also essential (Langston *et al.*, 2017; Masselink *et al.*, 2017). Some manipulative field experiments have been attempted, but logistical and even ethical issues mean these are uncommon (McKee and Mendelssohn, 1989; Tate and Battaglia, 2013; Abbott and Battaglia, 2015). Consequently, many studies employ controlled ‘flooding’ in greenhouse or ‘common garden’ experiments, although inevitably, experiments are constrained to focus on a limited species or habitat pool (van Zandt *et al.*, 2003; Hanley *et al.*, 2013, 2017, Li and Pennings, 2018). Many studies also impose long-term, or periodic, chronic salinity, rather than replicating the short-duration, acute immersion experienced immediately after a storm (Tolliver *et al.*, 1997; van Zandt and Mopper 2002; van Zandt *et al.*, 2003; Mopper *et al.*, 2016; Li and Pennings, 2018). A further problem is that rather than use natural seawater, experiments are often undertaken using commercially available

marine aquarium salt or even NaCl solutions (Sykes and Wilson 1988; Flynn *et al.*, 1995; Tolliver *et al.*, 1997; Mopper *et al.*, 2016), with no assessment of their validity as alternatives. In the second experiment described by **Hanley *et al.*, (2020a)** however, six different European sand dune plant species showed remarkable uniformity in stress and ecophysiological responses to marine aquarium salt versus locally collected seawater. This consistency suggests that the chemistry of the former is indeed close enough to the latter to use marine aquarium salt as a reliable experimental substitute.

Despite the various methodological problems, unsurprisingly perhaps, significant negative repercussions for plant survival, growth, and reproduction are apparent for plants subjected to seawater (or surrogate) immersion (van Zandt *et al.*, 2003; Mopper *et al.*, 2016; Hanley *et al.*, 2017, 2020a,b; Li and Pennings, 2018; Lum and Barton, 2020). Mortality is common, but even where plants survive short-pulses of seawater exposure subsequent recovery is compromised. A typical response to the ionic and osmotic shock associated with salinity is the accumulation of stress metabolites (e.g. proline) and ions (Ca^{2+} and K^{+}) to exclude or compartmentalize Na^{+} and Cl (Flowers and Colmer, 2008; Munns and Tester, 2008) (likely explaining why plant response to NaCl solution is more extreme than seawater which contains 1.2% Ca^{2+} and 1% K^{+}). Even if achieved however, a cost on plant fitness is probably inevitable (Munns and Tester, 2008; White *et al.*, 2014; Hanley *et al.*, 2020a,b).

Most importantly perhaps, the ability of plants to tolerate, and recover from, seawater flooding seems to be species-specific. Long-term observation of Arctic tundra following a major storm surge in the Mackenzie Delta, Canada, shows that dwarf shrub tundra had a

much-reduced regenerative capacity than graminoids or upright shrubs (Lantz *et al.*, 2015 - see also Middleton, 2009; Tate and Battaglia, 2013). Manipulative greenhouse experiments (Hanley *et al.*, 2017, 2020a; Li and Pennings, 2018; Edge *et al.*, 2020) generally corroborate field observations of species-specific variation. Working on two native Hawaiian plants, **Lum and Barton (2020)** for example, report not only species-specific variation in ecophysiological responses to increased salinity (imposed over 3-weeks), but also that tolerance increased for both species as plants aged. These observations represent a critical component of our understanding of plant response to the environmental pressures associated with SLR and storm surges. Not only is species-specific variation important, but it is essential to elucidate plant responses throughout ontogeny. Middleton (2009) for example describes species-specific variation in post-hurricane germination and recruitment ability of US Gulf Coast marshland species, a response ascribed principally to increased salinity. At the other end of the plant life cycle, **Hanley *et al.*, (2020b)** report how immersion of oilseed rape (*Brassica napus*) in seawater reduced seed yield, and perhaps most importantly, that growth of the resulting seedlings was also greatly reduced in comparison with progeny cultivated from non-flooded or even freshwater-flooded parent plants.

Although work in this area is anything but ‘mature’, these studies signpost flooding as a potential selective filter that could remove species from the post disturbance community. The loss of key species or functional groups from any vegetation is likely to compromise ecosystem processes and so limit the ability to supply essential ecosystem services. For vegetation like sand dunes, these losses may be particularly profound. In Florida for

example, Miller (2015) identified reduced cover of the dune building grass, *Uniola paniculata*, in low elevation areas subject to frequent flooding as a likely reason why dune erosion was more common in these sites. The interplay of ACC-linked changes in storm frequency and severity, with resulting shifts in plant community composition and thus resilience against further storm damage, is pivotal for understanding how ECEs contribute to coastal defence.

Inter-tidal Plant Communities

Communities subject to periodic, but predictable, (twice daily) tidal submersion and exposure to air – mangroves, saltmarshes and some algal communities.

Although mangrove forests are both a globally widespread and exceptionally important habitat for biodiversity and coastal defense provision in (sub)tropical regions, we focus here on the saltmarsh ecosystems more typically associated with temperate coastlines. This is simply because in this special issue, **Krauss and Ostler (2020)** provide a comprehensive review of how storms influence mangrove ecosystems and the vital ecosystem services they provide.

The physical damage caused by storms ranges from waves and strong currents dislodging or breaking above-ground tissue (Möller *et al.*, 2014), to complete denudation of vegetation (Morton and Barras, 2011). Fragmented or degraded marshes are generally more vulnerable to disturbance than intact habitat (Stagg *et al.*, 2020) and so are less resilient to extreme events. Responses also vary with vegetation height and stiffness (Vuik *et al.*, 2018). For example, when exposed to simulated storm conditions, the tall, rigid grass *Elymus athericus*

experienced more breakage than the shorter, more flexible *Puccinellia maritima* (Rupprecht *et al.*, 2017). Strong winds and water flows can tear the root mat from the marsh surface, laterally folding it into ridges – described by Cahoon (2006) as like ‘pushing a rug up along a wooden floor’. This alters marsh topography, lowering areas where turf was lost and raising elevations (up to 2 m) on the folded ridges (Guntenspergen *et al.*, 1995). This can affect long-term community recovery (Leonardi *et al.*, 2018; Mossman *et al.*, 2019).

In addition to direct damage, storms modify plant communities through changes to the physical environment (see reviews by Cahoon, 2006; Leonardi *et al.*, 2018). Storm-driven waves can cause lateral erosion of tidal flats and marshes (Callaghan *et al.*, 2010), with erosion of fronting tidal flats increasing marsh loss by amplifying the consistent pressure imposed by normal wind and wave action (Leonardi *et al.*, 2016). Saltmarshes are resistant to storm-driven erosion of the marsh surface however, with vegetation playing a key role in stabilizing the sediment (Spencer *et al.*, 2016). Importantly, significant amounts of sediment (mobilised from sub-tidal, intertidal or upstream areas) are deposited on saltmarshes during these events (de Groot *et al.*, 2011). For example, a single hurricane can deposit the equivalent of over a century of sediment accumulated in ‘normal’ conditions, and account for up to two thirds of long-term sedimentation (Williams and Flanagan, 2009). Burial under such rapid deposition can kill vegetation (Callaway and Zedler, 2004), and reduce growth and seedling establishment (Langlois *et al.*, 2001; Cao *et al.*, 2018). Marsh recovery following storm-driven sediment deposition can be rapid however, (Guntenspergen *et al.*, 1995) and increases in elevation improve colonization, particularly in subsiding marshes (Mendelssohn and Kuhn, 2003).

Storms can generate significant debris, either through breakage of local coastal vegetation or the remobilization of existing natural and artificial debris (Meixler, 2017). Like sediment, debris can kill or damage the vegetation beneath (Uhrin and Schellinger, 2011), modify environmental conditions such as sediment redox potential (Abbas *et al.*, 2014), and lead to reductions in species richness (Tate and Battaglia, 2013). The amount of damage depends on the type of debris deposited (Uhrin and Schellinger, 2011), the size of the mat and how long it persists (Valiela and Rietsma, 1995), so in some circumstances, recovery can be quick (Ehl *et al.*, 2017). Plant debris can also be important for propagule dispersal, but can act as a pathway for invasive species (Minchinton, 2006).

The impact of changes in soil salinity following storms is less clear. In some circumstances, high rainfall can ameliorate conditions, allowing plants to colonize or grow faster. For example, in the dry climate of California, Noe and Zedler (2001) found that heavy rainfall provided a window for germination by reducing soil salinity and increasing soil moisture.

Storms can also alter the inundation regime of tidal marshes through changes to coastal morphology that lead to closure of an estuary mouth or movements of tidal channels.

Zedler (2010) summarises how the storm-driven closure of the Tijuana estuary had substantial negative impacts on tidal marsh vegetation when subsequent drought caused moisture loss and hypersalinity in sediments.

More typical is the generally negative effect of seawater inundation; Janousek *et al.*, (2016) report how experimental increases in inundation over one growing season reduced plant productivity. It is also likely that even where tidal marsh plants survive storm disturbance, they are so ecophysiologicaly compromised that interactions with other species change.

The study by **Edge *et al.*, (2020)** on three European saltmarsh species is an excellent example. Following seawater immersion, the biomass of *Triglochin maritima* decreased markedly in mixed assemblages with *Plantago maritima* and *Aster tripolium*, compared to monoculture. Interestingly, *Plantago* performed markedly better in flooded, mixed assemblages than in monoculture, appearing to ‘take advantage’ of a relative decline in the growth of the other species (Hanley *et al.*, (2017) describe very similar shifts for supralittoral plants). Edge *et al.*, (2020) further note how that for 14 out of 18 trait-species combinations examined (including height, SLA, and leaf number), flooding response in mixed assemblages differed from monocultures, changing the direction, as well as magnitude, of flood effects. Plant trait and species composition shifts within saltmarsh communities are likely important to ecosystem stability and function (Ford *et al.*, 2016), but if disturbance associated with storm events facilitates the spread of non-native species, repercussions could be more severe. This is exactly what **Gallego-Tévar *et al.*, (2020)** report when they found that an invasive *Spartina* hybrid was better able to tolerate stressful post-flood salinity conditions than its parent species (see also Charbonneau *et al.*, 2017). Together, these studies underscore the importance of species identity in dictating community responses to storm disturbances, and thus the capacity of the saltmarsh ecosystem to continue to deliver key services as ACC continues.

Subtidal Plant Communities

Ecosystems continually submerged below sea-level – primarily seagrass beds, but includes marine macro-algal communities, most commonly kelp ‘forests’

Storm events can have substantial impacts on seagrass and macroalgal communities, from changes in the relative abundance of species within a community to total habitat loss. These impacts occur through physical disturbance from violent storms, burial by displaced sediment, and even subsequent ‘knock-on’ effects from pluvial flooding.

High wave energy and flow speeds can physically damage fronds and stipes (Denny *et al.*, 1989), uproot individuals (Preen *et al.*, 1995) or cause failure of holdfasts (Seymour *et al.*, 1989). While the biomechanics of storm effects are well understood (see Denny and Gaylord, 2002), predicting the impact of storm events is more complex. Structural damage and uprooting/ dislodgement can result in high mortality; for example, complete loss of giant kelp occurs in storm-intense years but is not seen everywhere (Edwards, 2004). Large, frequent and breaking waves exert the greatest forces and are most likely to result in structural damage or dislodgement, particularly in shallow water when a storm coincides with low tide (Preen *et al.*, 1995; Filbee-Dexter and Scheibling, 2012). Even moderate waves can lead to entanglement of kelp fronds, increasing the potential for tissue damage (Seymour *et al.*, 1989). Effects can vary according to substrate type, as wave-carried rocks can dislodge individuals, while sand grains and small pebbles scour roots and holdfasts or damage tissue (Shanks and Wright, 1986). Substrate type also affects the forces needed to dislodge macroalgae (Thomsen *et al.*, 2004).

Storm-driven waves do not affect every organism equally however. Vulnerability varies with spatial arrangement and age; individuals in the centre of algal stands are less likely to be removed by waves or strong currents, and small, young kelp are more easily dislodged than older, larger individuals (Thomsen *et al.*, 2004). Nonetheless, the higher biomass of

very large kelp makes them more susceptible to high wave energies (Seymour *et al.*, 1989). Consequently, severe storms can result in homogenization of age structure in kelp beds. Ecotypes or morphological plasticity provide resistance to high wave action (e.g. in shallow waters) (Fowler-Walker *et al.*, 2006), allowing some individuals or populations to better cope with an extreme event. Storms are also generally most frequent at the point in the annual cycle where organisms are most resistant (Burnett and Koehl, 2019); accordingly, changes to storm seasonality may have significant consequences for these communities.

In addition to the effects of wave action and shear stress, storm-generated waves and currents redistribute sediments, causing erosion in some areas and burial in others. Cabaco *et al.*, (2008) identified significant species-specific variation in seagrass tolerance to both burial with sediment and erosion. Recovery is generally rapid under shallow burial, but this capacity decreases markedly when more sediment is deposited (Fourqurean and Rutten, 2004; Gera *et al.*, 2014). Consequently, burial by up to 45 cm of sediment, reported following some severe storms (Kosciuch *et al.*, 2018; Browning *et al.*, 2019), is likely to lead to localized loss of communities.

As well as the impacts of storms at sea, heavy rainfall can have major impacts on sub-tidal ECEs via the discharge of nutrient-rich, sediment-laden freshwaters into coastal areas. These enriched waters cause turbidity and stimulate algal blooms and epiphytic growth, both of which lower light availability (Lapointe *et al.*, 2019). Seagrasses are especially vulnerable (Cobaco *et al.*, 2008), and impacts of flood-induced light limitation can be more severe than the physical impacts of storms (Carlson *et al.*, 2010). In addition, heavy rainfall can reduce salinity, particularly in lagoons or estuaries, sometimes for several months

(Herbeck *et al.*, 2011; Kowalski *et al.*, 2018,). Some seagrasses are intolerant of hyposaline conditions, leading to mortality and sub-lethal effects (Fernandez-Torquemada and Sanchez-Lizaso, 2011). Ridler *et al.*, (2006) observed that while thinning and leaf loss occurred immediately after hurricanes, further declines continued for many months likely due to low and fluctuating salinity. Tolerance to hyposalinity is however, variable between and within species, ecotype (Benjamin *et al.*, 1999) and season (Fernandez-Torquemada and Sanchez-Lizaso, 2011) reducing the predictability of how seagrass communities respond.

Storms are nonetheless important disturbance agents, and seagrasses can rapidly regrow from roots or rhizomes, despite substantial above-ground loss (Valiela *et al.*, 1998). Other macroalgae can reattach or regenerate when broken or dislodged (Thomsen and Wernberg, 2005). Furthermore, storms may actually facilitate medium and long distance dispersal of seagrass and macroalgae propagules (Bell *et al.*, 2008; Waters *et al.*, 2018) and be important in maintaining food web complexity, although increasing storm frequencies can challenge the ability of kelps to regrow and simplify food web structure (Byrnes *et al.*, 2011). Damage to kelp fronds can for example, stimulate grazing activity, so increasing potential tissue loss to an already stressed individual (O'Brien *et al.*, 2015). Reductions in canopy-forming macroalgae and seagrasses through a combination of direct storm damage and herbivory can lead to community shifts to opportunistic species, such as turf-forming algae (O'Brien *et al.*, 2015, Filbee-Dexter and Wernberg, 2018). Gaps resulting from the storm-driven loss of corals and other benthic animals can nevertheless facilitate macroalgal

423 colonization, particularly in the absence or reduction of herbivory (Edmunds, 2019;
424 Steneck *et al.*, 2019).

425 The impacts of extreme storm events are not experienced in isolation. Long-term
426 environmental changes, such as SLR, eutrophication and overfishing, influence community
427 susceptibility, as does the legacy of previous storms (i.e. position in the ‘storm recovery
428 cycle’). For example, substantial seagrass losses in North Queensland, Australia, were the
429 cumulative result of a succession of intense storm and flood years, urbanization, and
430 agricultural run-off, rather than the consequence of a single storm (McKenna *et al.*, 2015).
431 Storm events are also stressing systems already impacted by ACC, a combination that
432 could lead to higher losses than imposed by either driver in isolation (Babcock *et al.*, 2019).
433 Smale and Vance, (2016) for example report that while the cold-water kelp *Laminaria*
434 *hyperborea* was relatively resistant to storms, mixed stands containing warm water species,
435 such as *L. ochrolueca*, were more vulnerable. Consequently, observed and projected shifts
436 in kelp community composition due to increasing temperatures (Pessarrodona *et al.*, 2018)
437 could lead to greater kelp community vulnerability.

438 Collectively, the processes described above underpin observations of highly variable storm
439 impact on sub-tidal plant communities (Edwards, 2004; Filbee-Dexter and Scheibling,
440 2012). Long term studies can help identify the relative impacts of storms and anthropogenic
441 factors (Cuvillier *et al.*, 2017), but our understanding of storms on subtidal ECEs is limited
442 by few long term studies outside of coral reefs (Duffy *et al.*, 2019). While there are many
443 estimates of the impacts of single storms, it is rarely possible to put the patch-scale losses in
444 the context of the dynamics of the system. Despite advances with remote-sensing

techniques, the depth and turbidity of these systems mean that ground based observation will continue to be essential.

PLANT COMMUNITIES AND COASTAL DEFENCE

In addition to biodiversity loss, recent concern about the various threats to ECEs stems from their role in protecting agricultural land and urban communities from storm damage. Consequently, there is increasing focus on quantifying and valuing benefits associated with the ecosystem services provided by ECEs (Barbier *et al.* 2011, 2015; Temmerman *et al.*, 2013; Morris *et al.*, 2018). Although the methods used to generate accurate, global, economic estimates remain in their infancy (Barbier 2016), Costanza *et al.*, (2014) estimated that for tidal marshes alone, the provision of nursery grounds for commercial fisheries, carbon storage, recreation and flood protection provided US\$24.8 trillion to the global economy.

ECEs provide storm protection principally through the stabilization of substrates, and therefore the prevention of erosion, and attenuation of wave energy, and thus flood risk (Barbier 2015). Unlike hard (engineered) defences they are also dynamic; indeed the IPCC (2019) recognized how saltmarshes and mangroves can keep pace with fast rates of SLR ($> 10\text{mm yr}^{-1}$), depending on local variation in wave exposure, tidal range, sediment dynamics, and coastal land-use. Moreover, it is even possible that the extent of coastal wetlands (saltmarsh, freshwater marsh and mangrove) could increase by up to 60% because of SLR (Schuerch *et al.*, 2018). With appropriate management, supra-littoral sand dunes are also capable of adapting to shifts in sea levels and storm frequencies (Hanley *et al.*, 2014).

The growing evidence that ECEs reduce storm damage underpins their recognition as nature-based flood protection (Temmerman *et al.*, 2013; Narayan *et al.*, 2016; Van Coppenolle & Temmerman, 2019). The traditional approach to coastal defence has been to counter flood risk with ‘hard’ engineering, but measures like seawalls are expensive (up to £5,000 per m [Hudson *et al.*, 2015]), inflexible, and often deliver unexpected environmental outcomes (Firth *et al.*, 2014). Vegetated shorelines by contrast, are a natural defence and offer adaptability, flexibility and cost-effectiveness (e.g. £20 per m for dune stabilization (Hudson *et al.*, 2015)), with the additional benefit of the other ecosystem services they provide (Costanza *et al.*, 2014; Barbier 2015).

Protective role played by different ECEs

The protective value differs not only between ECEs, but also with regional and local geographical context. The principal defensive role played by dunes for example, stems from being a physical barrier to marine flooding, but their importance in this regard depends on local coastal geomorphology (e.g. sediment supply, land relief) and on the use and asset value of the land they protect (Hanley *et al.*, 2014). Dune vegetation stabilises substrates and reduces wave-driven erosion, with plant shoots reducing wave swash and roots increasing mechanical strength of the sediment (Feagin *et al.*, 2019), but even the identity of component species can be important. **de Battisti and Griffin (2020)** for example examined how three common European foredune species (*Ammophila arenaria*, *Cakile maritima*, and *Salsola kali*) varied in their ability to withstand simulated wave swash. Although *Ammophilla* was by far the most robust, by virtue of the protection provided by their roots, rhizomes and below ground shoots, all three species had a

488 remarkable capacity to tolerate wave action, underscoring how different plant species can
489 contribute to sand dune stability. (See also Charbonneau *et al.*, (2017) who report how
490 North American dunes stabilized by the invasive *Carex kobomugi* were less affected by
491 storm damage than those colonized by native *Ammophila breviligulata*). Nonetheless, de
492 Battisti and Griffin (2020) also show that despite an exceptionally well-developed
493 belowground shoot system, *Ammophila* resistance varies depending on sand particle size;
494 the coarser sediments associated with restored habitats increasing erosion potential
495 compared to finer sediment of natural regeneration sites. This finding is important since it
496 underscores why elucidation of biological and environmental factors is crucial to the
497 integration of natural habitats like sand dunes into coastal protection schemes. For other
498 supra-littoral habitats however, we understand little about their putative role in coastal
499 defence. Nonetheless, there is little doubt that coastal forests and freshwater wetlands
500 provide other vital ecosystem services like carbon sequestration and storage (see Stagg *et*
501 *al.*, 2020; Ury *et al.*, 2020).

502 The ability to track SLR (Kirwan *et al.*, 2016; IPCC, 2019) along with their well-known
503 capacity for wave attenuation (Möller *et al.*, 2014; Rupprecht *et al.*, 2017), has put
504 saltmarshes at the centre of current interest in ‘nature-based’ coastal defence solutions.
505 How effective wave attenuation is, depends strongly on topography (even to the extent of
506 friction imposed by the biogeomorphic landscape created by the plants) and (ontogenetic,
507 seasonal or species-specific) plant traits like shoot stiffness and density (Bouma *et al.*,
508 2010, 2014; Möller *et al.*, 2014). As a result, studies such as **Zhu *et al.* (2020)**, describing
509 variation in stem flexibility and breakability for a variety of European saltmarsh species,

are vital to understanding how communities will respond to increased storminess. Plant response can vary with wave conditions however. **Shao *et al.* (2020)** exposed *Spartina alterniflora* to different wave environments for 8 weeks and showed that key physiological and biochemical plant parameters varied accordingly; i.e. higher and more frequent waves imposed more stress. Nonetheless, wave-exposed plants tended to allocate more biomass to their roots, a response that may facilitate anchorage against wave impact. These biomechanical and morphological properties are likely to vary with plant age. **Cao *et al.*, (2020)** for instance describe how after seven weeks of simulated wave exposure, seedling survival and growth declined for all three common marshland species examined (*Spartina anglica*, *Scirpus maritimus* and *Phragmites australis*). Taken together these studies increase our understanding and prediction of spatio-temporal variation in saltmarsh community response to wave exposure, an essential pre-requisite in the design and implementation of nature-based flood protection.

In addition to species identity, age and seasonality, other marsh-specific characteristics are important determinants of wave attenuation. One of the key attributes is habitat size (Shepard *et al.*, 2011). Indeed, in a recent analysis of the long-term marsh persistence around the UK, Ladd *et al.*, (2019), revealed that marsh width was positively associated with higher sediment supply, although they noted also that current global declines in sediment flux are likely to diminish saltmarsh resilience to SLR. Although challenging, understanding the shifting dynamics of these regional-scale coastal processes is crucial to our ability to integrate marshes into coastal defence schemes (Bouma *et al.*, 2014, 2016). Not only is that because we need to know where and how ECEs fit into an integrated

coastal management approach, but long-term salt marsh persistence depends on continual recruitment of new plants.

For saltmarshes, propagule establishment often occurs on leading edges when sediment accretes on the adjacent ‘tidal flat’ (Bouma *et al.*, 2016). Even an apparently minor change in sediment levels may be sufficient to facilitate seedling establishment; an effect demonstrated by **Fivash *et al.*, (2020)** in their mesocosm experiment with the pioneer *Salicornia procumbens*. They show that elevation of sediment micro-topography by just 2 cm was the overwhelming driver of seedling growth (i.e. an average 25 % increase). They ascribed this response primarily to the effects of the ‘tidally driven oxygen pump’, i.e. increased emersion time allows more aeration of the raised sediment (see also Mossman *et al.*, 2019). Once pioneers like *Salicornia* have established, the environment they create (wave attenuation, sediment trapping and enhanced drainage) facilitates subsequent colonisation by later successional species and so the marsh can expand seaward (Temmerman *et al.*, 2007). Storms also have the potential to increase the landward marsh area if the habitat can retreat and displace terrestrial habitats. In these circumstances, **Kotter and Gedan (2020)** demonstrate that saltmarsh is pre-primed to take advantage of this opportunity, reporting how seeds of halophytic species can disperse up to 15 m into northeast American coastal pine forest. They argue that although saltwater intrusion will limit forest regeneration, the soil seed bank can thus support continued landward migration of saltmarsh species.

Much of the recent interest in mangroves stems from their perceived mitigation of the 2004 Indian Ocean Tsunami on coastal settlements. While their actual contribution remains

questionable (Barbier 2015), nonetheless, a number of studies report that mangroves can lower wave heights and reduce water levels during storm surges (Das and Vincent, 2009; Armitage *et al.*, 2019) and that their removal leads to increased coastal erosion and damage (Granek and Ruttenberg 2007; Barbier 2015). Like saltmarsh therefore, mangroves are at the forefront of contemporary research into how ECEs help defend our coastlines (see Krauss and Osland, 2020). It is also noteworthy, that Alongi (2008) highlights how much mangroves offer protection against extreme events is strongly linked to intrinsic habitat characteristics (these include forest location and width, tree density and size, soil texture), but also the presence of other ECEs, such as coral reefs, seagrass beds, and dunes.

The case for a substantial protective role of sub-tidal ECEs remains less clear (although coral reefs are well studied and widely believed to play a major role – see Barbier 2015). It is known however, that seagrasses attenuate wave energy (Christianen *et al.*, 2013; Reidenbach and Thomas, 2018), and thus likely offer some coastal defence (Barbier *et al.*, 2011; Ondiviela *et al.*, 2014). Furthermore, the reduction in wave energy seagrasses provide can reduce the erosion experienced by adjacent tidal marsh systems (Carr *et al.*, 2018) and stabilise or even facilitate beach expansion (James *et al.*, 2019). Consequently, the dramatic global decline of seagrass habitat is of great concern and underscores recent calls for wider habitat protection (Cullen-Unsworth and Unsworth 2018). It is less clear whether sub-tidal macroalgal communities play any role in wave attenuation and therefore coastal protection, but a full review is provided in this special issue (see **Morris *et al.*, 2020**). In short, Morris *et al.*, (2020) note how only a limited number of studies have investigated coastal protection, and in their own study in Australia found that wave

576 attenuation by the kelp *Ecklonia radiata* was restricted to a small subset of the
577 environmental conditions sampled.

578 *Using ECEs in integrated coastal defence*

579 The implementation of ‘soft’ or natural flood defences depends on landscape context
580 (including the economic value of the land threatened by SLR, erosion, and storm damage)
581 and whether it is actually feasible and cost-effective to maintain or move defences (Hoggart
582 *et al.*, 2014). The ‘hold the line’ option has been traditionally met by the construction of
583 ‘hard’ defences (engineered solutions utilising concrete walls, rocky breakwaters, steel
584 piling, or stone gabions) but these are extremely expensive and have limited ecological
585 value. There is nonetheless considerable interest in how we might ‘soften’ structures using
586 design alterations (e.g. modification of surface topography) to increase biodiversity value
587 (Firth *et al.*, 2014). It is also recognised that vegetated foreshores reduce wave impact on
588 sea walls, such that a fronting saltmarsh provides sufficient additional defence to allow sea
589 wall height to be lowered, with substantial savings to capital and maintenance costs (Vuik
590 *et al.*, 2016). Where natural habitat is absent, it may be possible to create it using
591 management actions to stabilize or accrete sediment. For example, the combination of
592 beach nourishment, sand traps and planting can establish sand dunes to provide storm
593 protection to landward hard defences (Feagin *et al.*, 2015). At the landscape scale, the
594 strategic integration of hard engineered and soft natural defences may provide the only
595 realistic, cost-effective way to protect large sections of coastline.

It is imperative however, to ensure that where integrated management is planned, an engineered intervention does not detrimentally affect nearby ECEs. For example, hard defences can disrupt natural coastal processes and sediment supply (Hanley *et al.*, 2014), while the problem of ‘coastal squeeze’ means that existing (or planned) ECEs fronting hard-engineered defences cannot always track SLR (Schuerch *et al.*, 2018). In these situations, the long-term sustainability of natural flood protection may be greater if there is the potential to move the line of defence landward. This can simply involve ensuring a capacity for an existing ECE to ‘roll back’ (see Kotter and Gedan, 2020), but increasingly, ECEs are created in former terrestrial habitats; a process often termed ‘managed retreat’ or ‘managed realignment’ (MR).

The most common example is the breaching of sea walls or dykes to allow tidal flooding with the expectation that newly inundated land will develop into saltmarsh. These schemes have met with mixed success however, many studies showing that the plant communities developing in MR sites differ from those in adjacent natural marshes (Mossman *et al.*, 2012; Masselink *et al.*, 2017). Environmental conditions, such as elevation in the tidal frame or geomorphic setting (Mossman *et al.*, 2012; Masselink *et al.*, 2017) are critical to successful restoration, but these alone are insufficient to explain all observed differences (Sullivan *et al.*, 2018). Propagule dispersal is often limited and limiting (Mossman *et al.*, 2012) and species-specific differences in dispersal ability could mean that early colonisers inhibit the establishment of later arriving species (Sullivan *et al.*, 2018). Planting species with low recruitment potential into newly established marshes could resolve this (Mossman *et al.*, 2019). A relative lack of topographic heterogeneity in MR sites may also limit

transition to saltmarsh (Masselink *et al.*, 2017; Lawrence *et al.*, 2018). As we have seen (Mossman *et al.*, 2019; Fivash *et al.*, 2020), even minor changes in surface elevation can have a substantial impact on seedling recruitment in saltmarsh. These studies highlight that, while MR often fails to deliver ‘natural’ saltmarshes, there is considerable potential for research-led management to improve restoration success.

SYNTHESIS AND FUTURE STUDIES

Although considerable research effort is focused on the response of ECEs to disturbance events, there remains both a geographical bias towards the US Gulf and Atlantic seaboard states, and limited understanding of how the multiple stressors associated with SLR, extreme storms, and other anthropogenic activities affect even a fraction of ECE species or habitats. Beyond a simplistic call for ‘more research with additional species and regions’, we discuss how illumination of plant species and community responses to flooding, sediment movement, mechanical damage and landscape-scale processes is needed to better inform our ability to manage the biodiversity of ECEs and ensure their continued contribution to coastal defence (Fig 1).

Research Priority I – *Effects of storm damage and flooding on plant reproductive performance and recruitment*

Parmesan and Hanley (2015) highlighted how despite a wealth of information detailing plant species and community response to the warming, drought and elevated atmospheric CO₂ (eCO₂) associated with ACC, remarkably little is known about how any of these factors influence plant regeneration biology. The same failing is true of ECE response to SLR and

storms, even though recruitment success is manifestly pivotal to understanding how environmental stress and perturbation influence plant community recovery. Indeed, it is at this point worth stressing that the disturbance associated with storms is an important, positive, factor in ECE dynamics. It is for example, well understood that tropical cyclones stimulate reproduction and open regeneration opportunities (Zimmerman *et al.*, 2018; Krauss and Osland 2020), while disturbance of sand dune vegetation is a key driver of plant biodiversity in these most dynamic of ecosystems (Green and Miller, 2019). What is less clear however, is how ACC-linked shifts in storm intensity and return times disrupt recruitment processes that have evolved in response to environmental dynamics typical of pre-industrial times (Hanley *et al.*, 2014; Imbert 2018).

Some experiments have focused on the effect of elevated salinity on flowering and reproduction, but all too often consider only long-term, chronic effects (e.g. Van Zandt and Mopper, 2002; Pathikonda *et al.*, 2010; Rajaniemi and Barrett, 2018). Nonetheless, these studies are important as they show; (a) responses may only become apparent long after exposure (Van Zandt and Mopper, 2002), (b) reduced sexual reproduction was not compensated by vegetative reproduction (Pathikonda *et al.*, 2010), and (c) germination potential is species-specific (Rajaniemi and Barrett, 2018). Many fewer authors report the impact of acute seawater flooding on the reproductive potential of coastal plants, but those that do evidence reduced flowering (White *et al.*, 2014; Hanley *et al.*, 2020a), and reproductive output (Hanley *et al.*, 2020b). A critical element of the latter study was that the growth of seedlings cultivated from parent plants subject to acute seawater immersion declined; i.e. while the parent plant might survive long enough to reproduce, longer-term

regeneration potential is compromised. The importance of changes in wave action on the dynamic sediment environment in saltmarsh regeneration may be better understood (Boauma *et al.*, 2016; Cao *et al.*, 2018), but there is a need to elucidate the effects of all manifestations of storm damage and flooding on plant reproductive and recruitment potential, including storm-driven dispersal.

Research Priority II –*Coastal plant responses to multiple-stressors associated with SLR and storm damage*

Teasing apart the interactive effects of saltwater flooding, mechanical damage, litter accumulation, and sediment shift on the plant community is challenging, a problem made all the more difficult simply because so few studies (outside the SE USA at least) have systematically examined how these different factors affect and shape plant community responses in isolation, let alone combination. Using remote imaging, Hauser *et al.*, (2015) report how saline inundation following Hurricane Sandy caused widespread wetland degradation in New Jersey, first by marsh dieback, and as a consequence, subsequent sediment erosion and retreat of the marsh inland. They also note the importance of plant community composition in this interaction; woody plants being more tolerant than herbaceous vegetation. Using an experimental approach, Tate and Battaglia (2013) considered the combined effects of seawater flooding and litter deposition. The application of locally sourced litter (degraded stems of black needlerush - *Juncus roemerianus*) to four plant communities along a Floridian estuarine gradient (brackish marsh, freshwater marsh, wetland forest, and pine savanna) had a profound negative effect on plant survival and species richness in all communities. In tandem with controlled seawater flooding however,

litter had a major impact on species composition in pine savannah, as salt-tolerant species capable of vegetative regrowth through dense detritus were the only species to persist. Tate and Battaglia (2013) also noted how vegetation in habitats with higher ambient sediment salinity was more resilient to the combined effects of flooding and litter deposition.

These studies (see also Imbert, 2018; Kendrick *et al.*, 2019) signpost the importance of interactive factors on the recovery of ECEs following storm and other ACC-linked disturbance events. Given the logistical issues associated with simultaneous replication or observation of multiple-stressors, it is unreasonable to expect a flurry of research focused on the interactive impacts of various storm disturbances on ECEs. Moreover, one could also argue that a true picture of coastal plant response needs also to consider eCO₂ and shifts in temperature and precipitation (Parmesan and Hanley, 2015). Indeed, Huang *et al.*, (2018) argued that an increase in night-time temperatures had facilitated the expansion of the shrub *Morella cerifera* into Virginian coastal grasslands with likely concomitant impacts on erosion regimes. Although by definition, unpicking the simultaneous interplay of several ACC-linked stressors is complex, as a first step studies could examine the responses of the same species to different stressors in isolation, and elucidate how at least two factors conspire to affect plant performance.

Research Priority III –*Plant community interactions and post-disturbance recovery*

Although it is well known that environmental perturbations (e.g. fire, herbivory, etc.), mediate plant community interactions, beyond a reasonable understanding of the role of tropical cyclones in forest dynamics (Hogan *et al.*, 2016; but see Pruitt *et al.*, 2019), the

704 impact of storms and SLR on plant-plant, plant-animal, and plant-microbial interactions in
705 ECEs is poorly resolved. We have discussed already how species-specific variation in plant
706 response to storms might act as a selective filter, removing susceptible species from the
707 recovering plant community. This is why field and multi-species (microcosm) greenhouse
708 experiments are invaluable; as shown by Hanley *et al.*, (2017) and Edge *et al.*, (2020), it is
709 by no means certain that plant species responses in monoculture are replicated in mixed
710 assemblages. Nonetheless, these kinds of study are rare and yet required to disentangle how
711 plant-plant interactions vary in response to a variety of storm-related impacts.

712 It is also worth stressing, that community interactions go beyond shifts in plant competitive
713 hierarchies. For example, although Camprubi *et al.*, (2012) report how three of six
714 Mediterranean sand dune species suffered complete mortality within a week of exposure to
715 seawater, the remainder had delayed or greatly reduced mortality when grown in
716 association with the mycorrhizal fungi, *Glomus intradices*. Symbiotic mycorrhizal fungi are
717 well known for their importance to plant health and vigour (Smith and Read, 2008), but in
718 coastal vegetation like sand dunes, the association may be essential for survival (Koske *et*
719 *al.*, 2004). Unfortunately, the vast majority of work on how the plant-mycorrhizal
720 association affects plant response to salinity comes from agricultural systems (Evelin *et al.*,
721 2019) and consequently we know little about how microbial symbionts respond to storm-
722 linked disturbances in ECEs, or how they moderate plant responses in the post-event
723 community.

724 Seawater inundation is also likely to have major effects on the soil physico-chemical
725 environment upon which all organisms depend. A detailed assessment of soil structure and

chemistry is beyond the scope of this review, but in addition to reduced aeration, increasing ionic concentrations and exchange capacity likely affect the bioavailability of key mineral nutrients (Kadiri *et al.*, 2012). Saline flooding will affect also soil microbial and invertebrate communities, and consequently, the decomposition and nutrient-cycling services they provide (Sjøgaard *et al.*, 2018; Stagg *et al.*, 2018). Remarkably few studies however, consider the impact of acute flooding on soil biogeochemistry, nor how additional stresses like sediment movement and litter accumulation affect soil dwelling animal and microbial communities and the processes they deliver.

Aboveground interactions are no less important. In an elegant experiment where sods of Louisianan marshland vegetation were exposed over 2-years to saline flood treatments, with and without herbivory, Gough and Grace (1999) reported that species loss was fastest in seawater treatments when mammal herbivores were also present. Although the flooding treatment was designed to mimic SLR rather than acute flooding, this study nonetheless emphasises how, even if species can tolerate one stress (flooding), the imposition of a second (herbivory) may filter species from the ecosystem (see also Mopper *et al.*, 2004; Schile and Mopper, 2006). Taken together, these studies underscore how post-storm conditions can affect plant morphology and the expression of defence metabolites, change herbivore performance and selection preferences, and how in combination, some plants may be excluded from the post-disturbance community. We cannot hope to understand how extreme storm events influence ECEs without a much greater understanding of these interactions.

Research Priority IV – *Better prediction of where and how storm events and SLR impact ECEs and the delivery of essential ecosystem services.*

Although we know that storms are more likely to happen with more frequency and greater intensity, a major challenge in predicting and understanding how ECEs will respond is to be able to forecast and define the range of storm surge and SLR scenarios for any given location. To achieve this, plant biologists must collaborate with geomorphologists, who with their understanding of bathymetry, wave dynamics, sediment supply, landform, and the biomechanical properties of vegetation, can offer vital insight into which ECEs are most susceptible and how they are likely to be affected (see also Krauss and Ostler, 2020). It also true, that in order to deliver accurate flood risk predictions and mitigation scenarios, geomorphologists must consider the contribution of plant communities to coastal processes.

The concept and application of coastal flood risk frameworks (CRAF) in coastal management is relatively well developed, but the focus has tended to be on how vulnerability to flooding affects human society rather than ECEs (Hallegatte *et al.*, 2013; Reimann *et al.*, 2018; Viavattene *et al.*, 2018). Nonetheless, there is developing appreciation that CRAF can be used to identify ‘at risk’ ecosystems (especially those that offer some measure of flood protection), or parts of the coastline where flood risk might be mitigated by virtue of the protection afforded by natural vegetation. In one such example, Christie *et al.* (2018) use the CRAF approach to pinpoint ‘hot spot’ sections of the North Norfolk (England) coast at greatest flood risk, and identify likely direct and indirect impacts based on an understanding of local geomorphology and hydrodynamic forcing during floods. Of particular note in this study is the finding that flood impact could be

reduced by saltmarsh; i.e. CRAF allows us to identify one of the key ecosystem services provided by coastal vegetation (see also Torresan *et al.*, 2012).

Another modelling approach, more familiar to plant biologists and ecologists, are species distribution models (SDMs). These have been widely used to predict how the geographical distribution of plant populations will respond to ACC-linked changes in precipitation and temperature (see Mairal *et al.*, 2018; Rodríguez-Rodríguez *et al.*, 2019). As noted already however, the combination of SLR with additional climate-change drivers is a unique, but largely ignored, issue for ECEs. Nonetheless, Garner *et al.* (2015) attempt some comparative synthesis, using SDM for Californian coastal plant species. They predict that by the end of this century, SLR alone threatens 60 of the 88 species considered and that 10 could completely lose their existing habitat range (due to flooding and erosion) within the (24,000km²) study region. This compares with only four species where shifts in temperature and precipitation alone eliminate all currently suitable habitats. Indeed, unlike plants threatened by SLR, some species may even gain suitable habitat space under likely temperature and precipitation scenarios. Garner *et al.* (2015) stress however, that in order to develop robust predictive models for coastal species, a much better mechanistic understanding of vegetation responses to SLR, flooding and climate scenarios is needed.

One way to achieve that aim is by undertaking long-term monitoring of threatened ECEs. This allows us to ‘ground truth’ predictive models by ‘back casting’ how recent environmental changes have actually influenced plant communities. By virtue of access to the Carolina Vegetation Survey, Ury *et al.*, (2020) were able to monitor changes in coastal forest communities over the past two decades. They report how the growth of tree species

like *Acer rubrum*, *Juniperus virginiana*, *Pinus serotina*, *Taxodium distichum* and various *Quercus* species was considerably reduced in low elevation sites where high soil salt content evidenced recent increased seawater seepage. In so doing, it is then possible to track how chronic saltwater intrusion has influenced tree growth and shifts in community composition over a 7-13 year time scale, exactly the kind of data needed to validate predictive models and understand how vulnerable ECEs respond to SLR, and changing storm frequencies and intensities. Long-term ecological surveys are time consuming and labour intensive, and for large coastlines therefore, impractical over the decadal timeframes in which we expect significant geomorphological and ecological changes to occur. Nonetheless, the use of remote sensing techniques in combination with localised ‘ground-truthing’ (see Stagg *et al.*, 2020) offers an effective combination to monitor and predict coastal change. The fact that both Stagg *et al.*, (2020) and Ury *et al.* (2020) highlight how the ability of coastal forests to deliver key ecosystem services is likely compromised by seawater inundation presents the most compelling reason to undertake long-term monitoring and predictive modelling studies into the future.

Conclusions - ECEs in Perspective

The threats posed by the myriad factors associated with ACC and changing storm patterns are worthy of considerable attention, not only from the many geomorphologists, environmental agencies and land managers already concerned with coastal defence, but also from biologists with any interest in plant ecophysiology or community ecology. Beyond any esoteric concern, as sea levels rise and the risk and impact of extreme storms increases, the associated economic repercussions will escalate. Hallegatte *et al.*, (2013) for example,

813 estimated that the costs associated with flooding for the 136 largest coastal cities would
814 increase from US\$6 billion in 2005, to US\$52 billion in 2050. Even under these extreme
815 circumstances, it seems unlikely that taxpayers will willingly subsidize the high cost of
816 protecting every vulnerable urban centre, transport link, or farm, with hard-engineered
817 defences. Given that coastal cities and food production globally are exposed to increasing
818 ACC-driven flood risk, nature-based risk mitigation, employing the conservation,
819 management, or even creation of ECEs with the capacity to track SLR and mitigate storm
820 surges seems ever more desirable. Indeed, the fact that Van Coppenolle & Temmerman
821 (2019) suggest how a cost-effective and dynamic answer (i.e. wetland creation) to the
822 problem of coastal defence can potentially be applied to over a third of the global land area
823 within the influence zone of storm surges, it would seem foolish to ignore the possibility.

824 A better understanding of the response of ECEs to seawater flooding, physical damage,
825 litter accumulation etc., at the levels of individual plant species (ecophysiological),
826 ecosystem (interactions), and landscape (distributions), can be delivered by plant scientists
827 from across our various disciplines. In turn, conservation biologists and ecologists can set
828 to work protecting and enhancing those habitats that deliver coastal defence. Only by so
829 doing can society hope to protect the unique biodiversity of our coastal habitats and the
830 essential ecosystem services they offer us in return.

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LITERATURE CITED

- Abbas A, Rubio-Casal A, de Cires A, et al. 2014.** Wrack burial reduces germination and establishment of the invasive cordgrass *Spartina densiflora*. *NeoBiota*, **21**: 65.
- Abbott MJ, Battaglia LL. 2015.** Purple pitcher plant (*Sarracenia rosea*) dieback and partial community disassembly following experimental storm surge in a coastal pitcher plant bog. *PLoS ONE* **10**: e0125475.
- Adame MF, Zaldivar-Jimenez A, Teutli C, et al. 2013.** Drivers of mangrove litterfall within a karstic region affected by frequent hurricanes. *Biotropica* **45**: 147– 154.
- Alongi DM. 2008.** Mangrove forests: resilience, protection from tsunamis, and responses to global climate change. *Estuarine, Coastal and Shelf Science* **76**: 1–13.
- Armitage AR, Weaver CA, Kominoski JS, Pennings SC. 2019.** Resistance to hurricane effects varies among vegetation types in the marsh-mangrove ecotone. *Estuaries and Coasts* doi.org/10.1007/s12237-019-00577-3.
- Babcock RC, Bustamante RH, Fulton EA, et al. 2019.** Severe continental-scale impacts of climate change are happening now: extreme climate events impact marine habitat forming communities along 45% of Australia's coast. *Frontiers in Marine Science* **6**: 14.
- Barbier EB 2015.** Valuing the storm protection service of estuarine and coastal ecosystems. *Ecosystem Services* **11**: 32-38.
- Barbier EB. 2016.** The protective value of estuarine and coastal ecosystem services in a wealth accounting framework. *Environmental and Resource Economics* **64**: 37-58.
- Barbier EB, Hacker SD, Kennedy C, et al. 2011.** The value of estuarine and coastal ecosystem services. *Ecological Monographs* **81**: 169-183.
- Barr JG, Engel V, Smith TJ, Fuentes JD. 2012.** Hurricane disturbance and recovery of energy balance, CO₂ fluxes and canopy structure in a mangrove forest of the Florida Everglades. *Agricultural and Forest Meteorology* **153**: 54– 66.
- Bell SS, Fonseca MS, Kenworthy WJ. 2008.** Dynamics of a subtropical seagrass landscape: links between disturbance and mobile seed banks. *Landscape Ecology* **23**: 67-74.

- 861 **Benjamin KJ, Walker DI, McComb AJ, Kuo J. 1999.** Structural response of marine and
862 estuarine plants of *Halophila ovalis* (R.Br.) Hook. f. to long-term hyposalinity.
863 *Aquatic Botany* **64**: 1-17.
- 864 **Bouma T, Vries MD, Herman PM. 2010.** Comparing ecosystem engineering efficiency of
865 two plant species with contrasting growth strategies. *Ecology* **91**: 2696–2704.
- 866 **Bouma TJ, van Belzen J, Balke T, et al. 2014.** Identifying knowledge gaps hampering
867 application of intertidal habitats in coastal protection: opportunities & steps to take.
868 *Coastal Engineering*, **87**: 147–157.
- 869 **Bouma TJ, van Belzen J, Balke T, et al.. 2016.** Short-term mudflat dynamics drive long-
870 term cyclic salt marsh dynamics. *Limnology and Oceanography* **61**: 2261-2275.
- 871 **Brown JK, Zinnert JC. 2018.** Mechanisms of surviving burial: Dune grass interspecific
872 differences drive resource allocation after sand deposition. *Ecography* **9**: e02162.
- 873 **Browning TN, Sawyer DE, Brooks GR, Larson RA, Ramos-Scharron CE, Canals-
874 Silander M. 2019.** Widespread deposition in a coastal bay following three major
875 2017 hurricanes (Irma, Jose, and Maria). *Scientific Reports* **9**: 13.
- 876 **Burnett NP, Koehl MAR. 2019.** Mechanical properties of the wave-swept kelp *Egregia
877 menziesii* change with season, growth rate and herbivore wounds. *Journal of
878 Experimental Biology* **222**: doi: 10.1242/jeb.190595.
- 879 **Byrnes JE, Reed DC, Cardinale BJ, Cavanaugh KC, Holbrook SJ, Schmitt RJ. 2011.**
880 Climate-driven increases in storm frequency simplify kelp forest food webs. *Global
881 Change Biology* **17**: 2513-2524.
- 882 **Cabaco S, Santos R, Duarte CM. 2008.** The impact of sediment burial and erosion on
883 seagrasses: A review. *Estuarine Coastal and Shelf Science* **79**: 354-366.
- 884 **Cahoon DR. 2006.** A review of major storm impacts on coastal wetland elevations.
885 *Estuaries and Coasts*, **29**: 889-898.
- 886 **Callaghan DP, Bouma TJ, Klaassen P, et al., 2010.** Hydrodynamic forcing on salt-marsh
887 development: Distinguishing the relative importance of waves and tidal flows.
888 *Estuarine, Coastal and Shelf Science*, **89**: 73-88.
- 889 **Callaway JC, Zedler JB. 2004.** Restoration of urban salt marshes: Lessons from southern
890 California. *Urban Ecosystems*, **7**: 107-124.

- 891 **Camprubi A, Abril M, Estaun V, Calvet C. 2012.** Contribution of arbuscular mycorrhizal
892 symbiosis to the survival of psammophilic plants after sea water flooding. *Plant and*
893 *Soil* **351**: 97–105.
- 894 **Carlson PR, Yarbrow LA, Kaufman KA, Mattson RA. 2010.** Vulnerability and resilience
895 of seagrasses to hurricane and runoff impacts along Florida's west coast.
896 *Hydrobiologia* **649**: 39-53.
- 897 **Carr J, Mariotti G, Fahgerazzi S, McGlathery K, Wiberg P. 2018.** Exploring the
898 impacts of seagrass on coupled marsh-tidal flat morphodynamics. *Frontiers of*
899 *Environmental Science* **6**: 92.
- 900 **Carter GA, Otvos EG, Anderson CP, Funderburk WR, Lucas KL. 2018.** Catastrophic
901 storm impact and gradual recovery on the Mississippi-Alabama barrier islands,
902 2005–2010: Changes in vegetated and total land area, and relationships of post-
903 storm ecological communities with surface elevation. *Geomorphology* **321**: 72-86.
- 904 **Castañeda-Moya E, Twilley RR, Rivera-Monroy VH, et al. 2010.** Sediment and nutrient
905 deposition associated with Hurricane Wilma in mangroves of the Florida Coastal
906 Everglades. *Estuaries and Coasts* **33**: 45– 58.
- 907 **Cao H, Zhu Z, Balke T, Zhang L, Bouma TJ. 2018.** Effects of sediment disturbance
908 regimes on *Spartina* seedling establishment: Implications for salt marsh creation and
909 restoration. *Limnology and Oceanography*, **63**: 647-659.
- 910 **Cao H, Zhu Z, James R, et al. 2020.** Wave effects on seedling establishment of three
911 pioneer marsh species: survival, morphology and biomechanics. *Annals of Botany*
912 **125**: doi.org/10.1093/aob/mcz136.
- 913 **Charbonneau BR, Wootton LS, Wnek JP, Langley JA, Posner MA. 2017.** A species
914 effect on storm erosion: Invasive sedge stabilized dunes more than native grass
915 during Hurricane Sandy. *Journal of Applied Ecology* **54**: 1–10.
- 916 **Christianen M.JA, van Belzen J, Herman PMJ, et al. 2013.** Low-canopy seagrass beds
917 still provide important coastal protection services. *PLoS One* **8**: e62413.
- 918 **Colmer TD, Voisenek LACJ. 2009.** Flooding tolerance: suites of plant traits in variable
919 environments. *Functional Plant Biology* **36**: 665-681.

- 920 **Costanza R, de Groot R, Sutton P, et al., 2014.** Changes in global value of ecosystem
921 services. *Global Environmental Change* **26**: 152-158.
- 922 **Cullen-Unsworth LC, Unsworth RFK. 2018.** A call for seagrass protection. *Science* **361**:
923 446–448.
- 924 **Cuvillier A, Villeneuve N, Cordier E, et al. 2016.** Causes of seasonal and decadal
925 variability in a tropical seagrass seascape (Reunion Island, South Western Indian
926 Ocean). *Estuarine, Coastal and Shelf Science* **184**: 90-101.
- 927 **Das S, Vincent JR. 2009.** Mangroves protected villages and reduced death toll during
928 Indian super cyclone. *Proceedings of the National Academy of Sciences* **106**: 7357-
929 7360.
- 930 **de Battisti D, Griffin JN. 2020.** Belowground biomass of plants, with a key contribution
931 of buried shoots, increases foredune resistance to wave swash. *Annals of Botany*
932 **125**: doi.org/10.1093/aob/mcz125
- 933 **de Groot AV, Veeneklaas RM, Bakker JP. 2011.** Sand in the salt marsh: Contribution of
934 high-energy conditions to salt-marsh accretion. *Marine Geology*, **282**: 240-254.
- 935 **Denny M, Brown V, Carrington E, Kraemer G, Miller A. 1989.** Fracture-mechanics and
936 the survival of wave-swept macroalgae. *Journal of Experimental Marine Biology*
937 *and Ecology* **127**: 211-228.
- 938 **Denny M, Gaylord B. 2002.** The mechanics of wave-swept algae. *Journal of Experimental*
939 *Biology* **205**: 1355-1362.
- 940 **Donguy J R, Meyers G. 1996.** Seasonal variations of sea surface salinity and temperature
941 in the tropical Indian Ocean. *Deep Sea Research Part I* **43**: 117–138.
- 942 **Douglas SH, Bernier JC, Smith KEL. 2018.** Analysis of multi-decadal wetland changes,
943 and cumulative impact of multiple storms 1984 to 2017. *Wetlands Ecology and*
944 *Management* **26**: 1121–1142.
- 945 **Doyle TW, Smith TJ, Robblee MB. 1995.** Wind damage effects of Hurricane Andrew on
946 mangrove communities along the southwest coast of Florida. *Journal of Coastal*
947 *Research* **21**: 159– 168.

- 948 **Duffy JE, Benedetti-Cecchi L, Trinanes J, et al. 2019.** Toward a coordinated global
 949 observing system for seagrasses and marine macroalgae. *Frontiers in Marine*
 950 *Science* **6**: doi: 10.3389/fmars.2019.00317.
- 951 **Edge RS, Sullivan MJP, Pedley SM, Mossman HL. 2020.** Species interactions modulate
 952 the response of saltmarsh plants to flooding. *Annals of Botany* **125**:
 953 doi.org/10.1093/aob/mcz120
- 954 **Edmunds PJ. 2019.** Three decades of degradation lead to diminished impacts of severe
 955 hurricanes on Caribbean reefs. *Ecology* **100**: e02587.
- 956 **Edwards MS. 2004.** Estimating scale-dependency in disturbance impacts: El Niños and
 957 giant kelp forests in the northeast Pacific. *Oecologia* **138**: 436-447.
- 958 **Ehl KM, Raciti SM, Williams JD. 2017.** Recovery of salt marsh vegetation after removal
 959 of storm-deposited anthropogenic debris: Lessons from volunteer clean-up efforts in
 960 Long Beach, NY. *Marine Pollution Bulletin*, **117**: 436-447.
- 961 **Evelin H, Devi TS, Gupta S, Kapoor R. 2019.** Mitigation of salinity stress in plants by
 962 arbuscular mycorrhizal symbiosis: Current understanding and new challenges.
 963 *Frontiers in Plant Science* **10**: 470.
- 964 **Feagin RA, Figlus J, Zinnert JC, et al. 2015.** Going with the flow or against the grain?
 965 The promise of vegetation for protecting beaches, dunes, and barrier islands from
 966 erosion. *Frontiers in Ecology and the Environment* **13**: 203– 210.
- 967 **Feagin RA, Furman M, Salgado K, et al. 2019.** The role of beach and sand dune
 968 vegetation in mediating wave run up erosion. *Estuarine, Coastal and Shelf Science*
 969 **219**: 97–106.
- 970 **Fernandez-Torquemada Y, Sanchez-Lizaso JL. 2011.** Responses of two Mediterranean
 971 seagrasses to experimental changes in salinity. *Hydrobiologia* **669**: 21-33.
- 972 **Filbee-Dexter K, Scheibling RE. 2012.** Hurricane-mediated defoliation of kelp beds and
 973 pulsed delivery of kelp detritus to offshore sedimentary habitats. *Marine Ecology*
 974 *Progress Series* **455**: 51-64.
- 975 **Filbee-Dexter K, Wernberg T. 2018.** Rise of turfs: a new battlefield for globally declining
 976 kelp forests. *Bioscience* **68**: 64-76.

- 977 **Firth LC, Thompson RC, Bohn K, et al. 2014.** Between a rock and a hard place:
 978 environmental and engineering considerations when designing coastal defence
 979 structures. *Coastal Engineering* **87**: 122-135.
- 980 **Fivash GS, Belzen JV, Temmink RJM, et al., 2020.** Micro-topography boosts growth
 981 rates in salt marsh pioneers by amplifying a tidally-driven oxygen pump:
 982 implications for restoration and recruitment. *Annals of Botany* **125**: doi:
 983 10.1093/aob/mcz137.
- 984 **Flowers TJ, Colmer TD. 2008.** Salinity tolerance in halophytes. *New Phytologist* **179**:
 985 945-963.
- 986 **Flynn KM, McKee KL, Mendelssohn IA. 1995.** Recovery of freshwater marsh vegetation
 987 after a saltwater intrusion event. *Oecologia* **103**: 63–72.
- 988 **Fourqurean JW, Rutten LM. 2004.** The impact of Hurricane Georges on soft-bottom,
 989 back reef communities: Site- and species-specific effects in south Florida seagrass
 990 beds. *Bulletin of Marine Science* **75**: 239-257.
- 991 **Fowler-Walker MJ, Wernberg T, Connell SD. 2006.** Differences in kelp morphology
 992 between wave sheltered and exposed localities: morphologically plastic or fixed
 993 traits? *Marine Biology* **148**: 755-767.
- 994 **Gallego-Tévar B, Grewell BJ, Futrell CJ, Drenovsky RE, Castillo JM. 2020.** Interactive
 995 effects of salinity and inundation on native *Spartina foliosa*, invasive *S. densiflora*,
 996 and their hybrid from San Francisco Estuary, California. *Annals of Botany* **125**:
 997 doi.org/10.1093/aob/mcz170.
- 998 **Garner KL, Chang MY, Fulda MT et al. 2015.** Impacts of sea level rise and climate
 999 change on coastal plant species in the central California coast. *PeerJ* **3**: e958.
- 1000 **Granek EF, Ruttenberg BI. 2007.** Protective capacity of mangroves during tropical
 1001 storms: a case study from ‘Wilma’ and ‘Gamma’ in Belize. *Marine Ecological*
 1002 *Progress Series* **343**: 101-105.
- 1003 **Gera A, Pages JF, Arthur R, Farina S, Roca G, Romero J, Alcoverro T. 2014.** The
 1004 effect of a centenary storm on the long-lived seagrass *Posidonia oceanica*.
 1005 *Limnology and Oceanography* **59**: 1910-1918.

- 1006 **Green MD, Miller TE. 2019.** Germination traits explain deterministic processes in the
1007 assembly of early successional coastal dune vegetation. *Estuaries and Coasts* **42**:
1008 1097–1103.
- 1009 **Gurbisz C, Kemp WM, Sanford LP, Orth RJ. 2016.** Mechanisms of storm-related loss
1010 and resilience in a large submersed plant bed. *Estuaries and Coasts* **39**: 951-966.
- 1011 **Guntenspergen GR, Cahoon DR, Grace J, et al. 1995.** Disturbance and recovery of the
1012 Louisiana coastal marsh landscape from the impacts of Hurricane Andrew. *Journal*
1013 *of Coastal Research*: 324-339.
- 1014 **Hallegatte S, Green C, Nicholls RJ, Corfee-Morlot J. 2013.** Future flood losses in major
1015 coastal cities. *Nature Climate Change* **3**: 802-806,
- 1016 **Haller WT, Sutton DL, Barlowe WC. 1974.** Effects of salinity on growth of several
1017 aquatic macrophytes. *Ecology* **55**: 891-894.
- 1018 **Hanley ME, Yip PYS, Hoggart SPG, et al. 2013.** Riding the storm: The response of
1019 *Plantago lanceolata* to simulated tidal flooding. *Journal of Coastal Conservation*
1020 **17**: 799-803.
- 1021 **Hanley ME, Hoggart SPG, Simmonds DJ, et al. 2014.** Shifting sands? Coastal protection
1022 by sand banks, beaches and dunes. *Coastal Engineering* **87**: 136-146.
- 1023 **Hanley ME, Gove TL, Cawthray GR, Colmer TD. 2017.** Differential responses of three
1024 coastal grassland species to seawater flooding. *Journal of Plant Ecology* **10**: 322–
1025 330.
- 1026 **Hanley ME, Sanders SKD, Stanton H-M, Billington RA, Boden R. 2020a.** A pinch of
1027 salt: Response of coastal grassland plants to simulated seawater inundation
1028 treatments. *Annals of Botany* **125**: doi 10.1093/aob/mcz042
- 1029 **Hanley ME, Hartley FC, Hayes L, Franco M. 2020b.** Simulated seawater flooding
1030 reduces oilseed rape growth, yield, and progeny performance. *Annals of Botany*
1031 **125**: doi.10.1093/aob/mcz026.
- 1032 **Harris AL, Zinnert JC, Young DR. 2017.** Differential response of barrier island dune
1033 grasses to species interactions and burial. *Plant Ecology* **218**: 609–619.

- 1034 **Hauser S, Meixler MS, Laba M. 2015.** Quantification of impacts and ecosystem services
1035 loss in New Jersey coastal wetlands due to Hurricane Sandy storm surge. *Wetlands*
1036 **35:** 1137-1148.
- 1037 **Herbeck LS, Unger D, Krumme U, Liu SM, Jennerjahn TC. 2011.** Typhoon-induced
1038 precipitation impact on nutrient and suspended matter dynamics of a tropical
1039 estuary affected by human activities in Hainan, China. *Estuarine Coastal and Shelf*
1040 *Science* **93:** 375-388.
- 1041 **Hogan JA, Zimmerman JK, Thompson J, Nytch CJ, Uriarte M. 2016.** The interaction
1042 of land-use legacies and hurricane disturbance in subtropical wet forest: twenty-one
1043 years of change. *Ecosphere* **7:** e01405
- 1044 **Hoggart SPG, Hanley ME, Parker DJ, et al. 2014.** The consequences of doing nothing:
1045 The effects of seawater flooding on coastal zones. *Coastal Engineering* **87:** 169-
1046 182.
- 1047 **Huang H, Zinnert JC, Wood LK, Young DR, D'Odorico P. 2018.** A non-linear shift
1048 from grassland to shrubland in temperate barrier islands. *Ecology* **99:** 1671-1681.
- 1049 **Hudson T, Keating K, Pettit A. 2015.** *Cost estimation for coastal protection – summary*
1050 *of evidence*. Environment Agency UK, Report no. SC080039/R7.
- 1051 **Imbert D. 2018.** Hurricane disturbance and forest dynamics in east Caribbean mangroves.
1052 *Ecosphere* **9:** e02231.
- 1053 **IPCC, 2019.** Summary for Policymakers. In: *IPCC Special Report on the Ocean and*
1054 *Cryosphere in a Changing Climate* [Pörtner H-O, Roberts DC, Masson-Delmotte V,
1055 P. et al., (eds.)]. IPCC, Geneva, Switzerland, In press.
- 1056 **James RK, Silva R, van Tussenbroek BI et al., 2019.** Maintaining tropical beaches with
1057 seagrass and algae: A promising alternative to engineering solutions. *BioScience* **69:**
1058 136–142.
- 1059 **Janousek CN, Buffington KJ, Thorne KM, Guntenspergen GR, Takekawa JY,**
1060 **Dugger BD. 2016.** Potential effects of sea-level rise on plant productivity: species-
1061 specific responses in northeast Pacific tidal marshes. *Marine Ecology Progress*
1062 *Series*, **548:** 111-125.

- 1063 **Kadiri M, Spencer KL, Heppell CM. 2012,** Potential contaminant release from
1064 agricultural soil and dredged sediment following managed realignment. *Journal of*
1065 *Soils and Sediments* **12**: 1581– 1592.
- 1066 **Kendrick GA, Nowicki RJ, Olsen YS, et al., 2019.** A systematic review of how multiple
1067 stressors from an extreme event drove ecosystem-wide loss of resilience in an iconic
1068 seagrass community. *Frontiers in Marine Science* **6**: 455.
- 1069 **Kirwan ML, Temmerman S, Skeeahan EE, Guntenspergen GR, Fagherazzi S. 2016.**
1070 Overestimation of marsh vulnerability to sea level rise. *Nature Climate Change* **6**:
1071 253– 260.
- 1072 **Kosciuch TJ, Pilarczyk JE, Hong I, et al. 2018.** Foraminifera reveal a shallow nearshore
1073 origin for overwash sediments deposited by Tropical Cyclone Pam in Vanuatu
1074 (South Pacific). *Marine Geology* **396**: 171-185.
- 1075 **Koske RE, Gemma JN, Corkidi L, Siguenza C, Rincon E. 2004.** Arbuscular
1076 mycorrhizas in coastal dunes. In: Martinez ML, Psuty NP. (eds.) *Coastal Dunes,*
1077 *Ecology and Conservation*, Springer, Heidelberg, pp. 173-187.
- 1078 **Kottler EJ, Gedan K. 2020.** Seeds of change: will the soil seed bank support marsh
1079 migration? *Annals of Botany* **125**: doi 10.1093/aob/mcz133
- 1080 **Kowalski JL, DeYoe HR, Boza GH, Hockaday DL, Zimba PV. 2018.** A comparison of
1081 salinity effects from Hurricanes Dolly (2008) and Alex (2010) in a Texas Lagoon
1082 System. *Journal of Coastal Research* **34**: 1429-1438.
- 1083 **Krauss KW Osland MJ. 2020.** Tropical cyclones and the organization of mangrove
1084 forests: a review. *Annals of Botany* **125**: doi.org/10.1093/aob/mcz161.
- 1085 **Ladd CJT, Duggan-Edwards MF, Bouma TJ, Pagès JF, Skov MW. 2019.** Sediment
1086 supply explains long-term and large-scale patterns in saltmarsh lateral expansion
1087 and erosion. *Geophysical Research Letters* doi.org/10.1029/2019GL083315.
- 1088 **Langlois E, Bonis A, Bouzillé JB. 2001.** The response of *Puccinellia maritima* to burial: A
1089 key to understanding its role in salt-marsh dynamics? *Journal of Vegetation*
1090 *Science*, **12**: 289-297.

- 1091 **Langston AK, Kaplan DA, Putz FE. 2017.** A casualty of climate change? Loss of
1092 freshwater forest islands on Florida's Gulf Coast. *Global Change Biology* **23**: 5383–
1093 5397.
- 1094 **Lantz TC, Kokelj SV, Fraser RH. 2015.** Ecological recovery in an Arctic delta following
1095 widespread saline incursion. *Ecological Applications* **25**: 172–185.
- 1096 **Lapointe BE, Herren LW, Brewton RA, Alderman P. 2019.** Nutrient over-enrichment
1097 and light limitation of seagrass communities in the Indian River Lagoon, an
1098 urbanized subtropical estuary. *Science of the Total Environment* **699**: 134068.
- 1099 **Larkum AWD, West RJ. 1990.** Long-term changes of seagrass meadows in Botany Bay,
1100 Australia. *Aquatic Botany* **37**: 55-70.
- 1101 **Lawrence PJ, Smith GR, Sullivan MJP, Mossman HL. 2018.** Restored saltmarshes lack
1102 the topographic diversity found in natural habitat. *Ecological Engineering* **115**: 58–
1103 66.
- 1104 **Leonardi N, Ganju NK, Fagherazzi S. 2016.** A linear relationship between wave power
1105 and erosion determines salt-marsh resilience to violent storms and hurricanes.
1106 *Proceedings of the National Academy of Sciences*, **113**: 64-68.
- 1107 **Leonardi N, Carnacina I, Donatelli C, et al., 2018.** Dynamic interactions between coastal
1108 storms and salt marshes: A review. *Geomorphology* **301**: 92–107.
- 1109 **Li F, Pennings SC. 2018.** Responses of tidal freshwater and brackish marsh macrophytes
1110 to pulses of saline water simulating sea level rise and reduced discharge. *Wetlands*
1111 **38**: 885–891.
- 1112 **Lum TD, Barton KE. 2020.** Ontogenetic variation in salinity tolerance and ecophysiology
1113 of coastal dune plants. *Annals of Botany* **125**: doi.org/10.1093/aob/mcz097.
- 1114 **Mairal M, Caujapé-Castells J, Pellissier L, et al. 2018.** A tale of two forests: ongoing
1115 aridification drives population decline and genetic diversity loss at continental scale
1116 in Afro-Macaronesian evergreen-forest archipelago endemics. *Annals of Botany*
1117 **122**: 1005– 1017.
- 1118 **Malloch AJC, Bamidele JF, Scott AM. 1985.** The phytosociology of British sea-cliff
1119 vegetation with special reference to the ecophysiology of some maritime cliff
1120 plants. *Vegetatio* **62**: 309-317.

- 1121 **Masselink G, Scott T, Poate T, et al. 2015.** The extreme 2013/14 winter storms:
 1122 Hydrodynamic forcing and coastal response along the southwest coast of England.
 1123 *Earth Surface Processes and Landforms* **41**: 378-391.
- 1124 **Masselink G, Hanley ME, Halwyn AC, et al., 2017.** Evaluation of salt marsh restoration
 1125 by means of self-regulating tidal gate – Avon estuary, south Devon, UK. *Ecological*
 1126 *Engineering* **106**: 174-190.
- 1127 **McKee KL, Mendelssohn IA 1989.** Response of a freshwater marsh plant community to
 1128 increased salinity and increased water level. *Aquatic Botany* **34**: 301-316.
- 1129 **McKenna S, Jarvis J, Sankey T, et al. 2015.** Declines of seagrasses in a tropical harbour,
 1130 North Queensland, Australia, are not the result of a single event. *Journal of*
 1131 *Biosciences* **40**: 389-398.
- 1132 **Meixler MS. 2017.** Assessment of Hurricane Sandy damage and resulting loss in
 1133 ecosystem services in a coastal-urban setting. *Ecosystem Services* **24**: 28-46
- 1134 **Mendelssohn IA, Kuhn NL. 2003.** Sediment subsidy: effects on soil-plant responses in a
 1135 rapidly submerging coastal salt marsh. *Ecological Engineering*, **21**: 115-128.
- 1136 **Middleton EA. 2009.** Regeneration of coastal marsh vegetation impacted by hurricanes
 1137 Katrina and Rita. *Wetlands* **29**: 54-65.
- 1138 **Millennium Ecosystem Assessment, 2005.** *Ecosystems and Human Well-being:*
 1139 *Biodiversity Synthesis*. World Resources Institute, Washington, DC.
- 1140 **Minchinton TE. 2006.** Rafting on wrack as a mode of dispersal for plants in coastal
 1141 marshes. *Aquatic Botany*, **84**: 372-376.
- 1142 **Möller I, Kudella M, Rupprecht F, et al. 2014.** Wave attenuation over coastal salt
 1143 marshes under storm surge conditions. *Nature Geoscience* **7**: 727-731.
- 1144 **Mopper S, Wang YY, Criner C, Hasenstein K. 2004.** *Iris hexagona* hormonal responses
 1145 to salinity stress, leafminer herbivory, and phenology. *Ecology* **85**: 38-47.
- 1146 **Mopper S, Wiens KC, Goranova GA. 2016.** Competition, salinity, and clonal growth in
 1147 native and introduced irises. *American Journal of Botany* **103**: 1575-1581.
- 1148 **Morris RL, Konlechner TM, Ghisalberti M, Swearer SE. 2018.** From grey to green:
 1149 Efficacy of eco-engineering solutions for nature-based coastal defence. *Global*
 1150 *Change Biology* **24**: 1827-1842.

- 1151 **Morris RL, Graham TDJ, Kelvin J, Ghisalberti M, Swearer SE. 2020.** Kelp beds as
 1152 coastal protection: wave attenuation of *Ecklonia radiata* in a shallow coastal bay.
 1153 *Annals of Botany* 125: doi.org/10.1093/aob/mcz127.
- 1154 **Morton RA, Barras JA. 2011.** Hurricane impacts on coastal wetlands: A half-century
 1155 record of storm-generated features from southern Louisiana. *Journal of Coastal*
 1156 *Research*, **27**: 27-43.
- 1157 **Mossman HL, Davy AJ, Grant A. 2012.** Does managed coastal realignment create
 1158 saltmarshes with 'equivalent biological characteristics' to natural reference sites?
 1159 *Journal of Applied Ecology* **49**: 1446-1456.
- 1160 **Mossman HL, Grant A, Davy AJ. 2019.** Manipulating saltmarsh microtopography
 1161 modulates the effects of elevation on sediment redox potential and halophyte
 1162 distribution. *Journal of Ecology* doi.10.1111/1365-2745.13229.
- 1163 **Munns R, Tester M. 2008.** Mechanisms of salt tolerance. *Annual Review of Plant Biology*
 1164 **59**: 651-681.
- 1165 **Narayan S, Beck MW, Reguero BG, et al. 2016.** The effectiveness, costs and coastal
 1166 protection benefits of natural and nature-based defences. *PLoS ONE* **11**: e0154735.
- 1167 **Negrão S, Schmöckel SM, Tester M. 2017.** Evaluating physiological responses of plants
 1168 to salinity stress. *Annals of Botany* **119**: 1–11.
- 1169 **Noe GB, Zedler JB. 2001.** Variable rainfall limits the germination of upper intertidal
 1170 marsh plants in southern California. *Estuaries*, **24**: 30-40.
- 1171 **Noe G, Krauss K, Lockaby B, Conner WH, Hupp CR. 2013.** The effect of increasing
 1172 salinity and forest mortality on soil nitrogen and phosphorus mineralization in tidal
 1173 freshwater forested wetlands. *Biogeochemistry* 114: 225–244.
- 1174 **O'Brien JM, Scheibling RE, Krumhansl KA. 2015.** Positive feedback between large-
 1175 scale disturbance and density-dependent grazing decreases resilience of a kelp bed
 1176 ecosystem. *Marine Ecology Progress Series* **522**: 1-13.
- 1177 **Ondiviela B, Losada IJ, Lara JL, et al. 2014.** The role of seagrasses in coastal protection
 1178 in a changing climate. *Coastal Engineering* **87**: 158–168.

- 1179 **Paling EI, Kobryn HT, Humphreys G. 2008.** Assessing the extent of mangrove change
1180 caused by Cyclone Vance in the eastern Exmouth Gulf, northwestern Australia.
1181 *Estuarine, Coastal and Shelf Science* **77**: 603– 613.
- 1182 **Parmesan C, Hanley ME. 2015.** Plants and climate change: complexities and surprises.
1183 *Annals of Botany* **115**: 849-864.
- 1184 **Pathikonda S, Meerow A, He Z, Mopper S. 2010.** Salinity tolerance and genetic
1185 variability in freshwater and brackish *Iris hexagona* colonies. *American Journal of*
1186 *Botany* **97**: 1438-1443.
- 1187 **Pessarrodona A Foggo A, Smale DA. 2018.** Can ecosystem functioning be maintained
1188 despite climate-driven shifts in species composition? Insights from novel marine
1189 forests. *Journal of Ecology* **107**: 91– 104.
- 1190 **Platt WJ, Doren RF, Armentano, TV. 2000.** Effects of Hurricane Andrew on stands of
1191 slash pine (*Pinus elliotii* var. *densa*) in the everglades region of south Florida. *Plant*
1192 *Ecology* **146**: 43– 60.
- 1193 **Platt WJ, Joseph D Ellair DP. 2015.** Hurricane wrack generates landscape-level
1194 heterogeneity in coastal pine savanna. *Ecography* **38**: 63–73.
- 1195 **Preen AR, Long WJL, Coles RG. 1995.** Flood and cyclone related loss, and partial
1196 recovery, of more than 1000 km² of seagrass in Hervey-Bay, Queensland, Australia.
1197 *Aquatic Botany* **52**: 3-17.
- 1198 **Pruitt JN, Little AG, Majumdar SJ, Schoener TW, Fisher DN 2019.** Call-to-Action: A
1199 global consortium for tropical cyclone ecology. *Trends in Ecology and Evolution*
1200 doi.org/10.1016/j.tree.2019.04.009.
- 1201 **Rajaniemi TK, Barrett DT. 2018.** Germination responses to abiotic stress shape species
1202 distributions on coastal dunes. *Plant Ecology* **219**: 1271–1282.
- 1203 **Reidenbach MA, Thomas EL. 2018.** Influence of the seagrass, *Zostera marina*, on wave
1204 attenuation and bed shear stress within a shallow coastal bay. *Frontiers in Marine*
1205 *Science* **5**: 397.
- 1206 **Reimann L, Vafeidis AT, Brown S, Hinkel J, Tol RSJ. 2018.** Mediterranean UNESCO
1207 World Heritage at risk from coastal flooding and erosion due to sea-level rise.
1208 *Nature Communications* **9**: 4161.

- 1209 **Ridler MS, Dent RC, Arrington DA. 2006.** Effects of two hurricanes on *Syringodium*
1210 *filiforme*, manatee grass, within the Loxahatchee River Estuary, southeast Florida.
1211 *Estuaries and Coasts* **29**: 1019-1025.
- 1212 **Rodríguez-Rodríguez P, de Castro AGF, Seguí J, Traveset A, Sosa PA 2019.** Alpine
1213 species in dynamic insular ecosystems through time: conservation genetics and
1214 niche shift estimates of the endemic and vulnerable *Viola cheiranthifolia*. *Annals of*
1215 *Botany* **123**: 505-519.
- 1216 **Rupprecht F, Möller I, Paul M, et al. 2017.** Vegetation-wave interactions in salt marshes
1217 under storm surge conditions. *Ecological Engineering*, **100**: 301-315.
- 1218 **Sachithanandam V, Mageswaran T, Sridhar R, Purvaja R, Ramesh R. 2014.**
1219 Assessment of Cyclone Lehar's impact on seagrass meadows in Ross and Smith
1220 Islands, North Andaman. *Natural Hazards* **72**: 1253-1258.
- 1221 **Schile L, Mopper S. 2006.** The deleterious effects of salinity stress on leafminers and their
1222 freshwater host. *Ecological Entomology* **31**: 345– 351.
- 1223 **Schuerch M, Vafeidis A, Slawig T, Temmerman S. 2013.** Modelling the influence of
1224 changing storm patterns on the ability of a salt marsh to keep pace with sea level
1225 rise. *Journal of Geophysical Research: Earth Surface*, **118**: 84-96.
- 1226 **Schuerch S, Spencer T, Temmerman S, et al., 2018.** Future response of global coastal
1227 wetlands to sea-level rise. *Nature* **561**: 231–234.
- 1228 **Schwarz C, Brinkkemper J, Ruessink G. 2019.** Feedbacks between biotic and abiotic
1229 processes governing the development of foredune blowouts: A Review. *Journal of*
1230 *Marine Science and Engineering* **7**: doi.org/10.3390/jmse7010002
- 1231 **Seymour RJ, Tegner MJ, Dayton PK, Parnell PE. 1989.** Storm wave-induced mortality
1232 of giant-kelp, *Macrocystis pyrifera*, in southern-California. *Estuarine Coastal and*
1233 *Shelf Science* **28**: 277-292.
- 1234 **Shanks AL, Wright WG. 1986.** Adding teeth to wave action - the destructive effects of
1235 wave-borne rocks on intertidal organisms. *Oecologia* **69**: 420-428.
- 1236 **Shao D, Zhou W, Bouma TJ, et al., 2020.** Physiological and biochemical responses of the
1237 salt-marsh plant *Spartina alterniflora* to long-term wave exposure. *Annals of Botany*
1238 **125**: doi 10.1093/aob/mcz067

- 1239 **Shepard CC, Crain CM, Beck MW. 2011.** The protective role of coastal marshes: A
1240 systematic review and meta-analysis. *PLoS ONE* **6**: e27374.
- 1241 **Sjøgaard KS, Valdemarsen TB, Treusch AH. 2018.** Responses of an agricultural soil
1242 microbiome to flooding with seawater after managed coastal realignment.
1243 *Microorganisms* **6**: doi: 10.3390/microorganisms6010012.
- 1244 **Smale DA, Vance T. 2016.** Climate-driven shifts in species' distributions may exacerbate
1245 the impacts of storm disturbances on North-east Atlantic kelp forests. *Marine and*
1246 *Freshwater Research* **67**: 65-74.
- 1247 **Smith SE, Read DJ. 2008.** *Mycorrhizal Symbiosis*, 3rd edn, Elsevier, Academic Press, New
1248 York, USA.
- 1249 **Spencer T, Möller I, Rupprecht F, et al. 2016.** Salt marsh surface survives true-to-scale
1250 simulated storm surges. *Earth Surface Processes and Landforms*, **41**: 543-552.
- 1251 **Stagg CL, Baustian MM, Perry CL, Carruthers TJB, Hall CT. 2018.** Direct and
1252 indirect controls on organic matter decomposition in four coastal wetland
1253 communities along a landscape salinity gradient. *Journal of Ecology* **106**: 655-670.
- 1254 **Stagg CL, Osland MJ, Moon JA, et al., 2020.** Quantifying hydrologic controls on local-
1255 and landscape-scale indicators of coastal wetland loss. *Annals of Botany* **125**: doi
1256 10.1093/aob/mcz144.
- 1257 **Steinke TD, Ward CJ. 1989.** Some effects of the cyclones Domoina and Imboa on
1258 mangrove communities in the St. Lucia estuary, South Africa. *South African*
1259 *Journal of Botany* **55**: 340– 348.
- 1260 **Steneck RS, Arnold SN, Boenish R, et al. 2019.** Managing recovery resilience in coral
1261 reefs against climate-induced bleaching and hurricanes: a 15 year case study from
1262 Bonaire, Dutch Caribbean. *Frontiers in Marine Science* **6**: doi:
1263 10.3389/fmars.2019.00265.
- 1264 **Steyer GD, Perez BC, Piazza SC, Suir G 2007.** Potential consequences of saltwater
1265 intrusion associated with hurricanes Katrina and Rita. In: *Science and the storms –*
1266 *the USGS response to the hurricanes of 2005*. Report 13066C, Reston Virginia,
1267 USA.

- 1268 **Sullivan MJP, Davy AJ, Grant A, Mossman HL. 2018.** Is saltmarsh restoration success
1269 constrained by matching natural environments or altered succession? A test using
1270 niche models. *Journal of Applied Ecology* **55**: 1207– 1217.
- 1271 **Sykes MT, Wilson JB. 1988.** An experimental investigation into the response of some
1272 New Zealand sand dune species to salt spray. *Annals of Botany* **62**: 159-166.
- 1273 **Tate AS, Battaglia LL. 2013.** Community disassembly and reassembly following
1274 experimental storm surge and wrack application. *Journal of Vegetation Science* **24**:
1275 46-57.
- 1276 **Temmerman S, Bouma TJ, van de Koppel J, et al. 2007.** Vegetation causes channel
1277 erosion in a tidal landscape. *Geology* **35**: 631–634.
- 1278 **Temmerman S, Govers G, Wartel S, Meire P. 2004.** Modelling estuarine variations in
1279 tidal marsh sedimentation: Response to changing sea level and suspended sediment
1280 concentrations. *Marine Geology* **212**: 1– 19.
- 1281 **Temmerman S, Meire P, Bouma TJ, et al. 2013.** Ecosystem-based coastal defence in the
1282 face of global change. *Nature* **504**: 79–83.
- 1283 **Thomsen MS, Wernberg T. 2005.** Miniview: What affects the forces required to break or
1284 dislodge macroalgae? *European Journal of Phycology* **40**: 139-148.
- 1285 **Thomsen MS, Wernberg T, Kendrick GA. 2004.** The effect of thallus size, life stage,
1286 aggregation, wave exposure and substratum conditions on the forces required to
1287 break or dislodge the small kelp *Ecklonia radiata*. *Botanica Marina* **47**: 454-460.
- 1288 **Tolliver KS, Martin DW, Young DR. 1997.** Freshwater and saltwater flooding response
1289 for woody species common to barrier island swales. *Wetlands* **17**: 10-18.
- 1290 **Torresan S, Critto A, Rizzi J, Marcomini A. 2012.** Assessment of coastal vulnerability to
1291 climate change hazards at the regional scale: the case study of the north Adriatic
1292 sea. *Natural Hazards and Earth System Sciences* **12**: 2347-2368.
- 1293 **Uhrin AV, Schellinger J. 2011.** Marine debris impacts to a tidal fringing-marsh in North
1294 Carolina. *Marine Pollution Bulletin*, **62**: 2605-2610.
- 1295 **Ury EA, Anderson SM, Peet RK, Bernhardt ES, Wright JP. 2020.** Succession,
1296 regression and loss: does evidence of saltwater exposure explain recent changes in

- 1297 the tree communities of North Carolina's Coastal Plain? *Annals of Botany* **125**:
 1298 doi.org/10.1093/aob/mcz039.
- 1299 **Valiela I, Rietsma CS. 1995.** Disturbance of salt marsh vegetation by wrack mats in Great
 1300 Sippewissett Marsh. *Oecologia*, **102**: 106-112.
- 1301 **Valiela I, Peckol P, D'Avanzo C, et al. 1998.** Ecological effects of major storms on
 1302 coastal watersheds and coastal waters: Hurricane Bob on Cape Cod. *Journal of*
 1303 *Coastal Research* **14**: 218-238.
- 1304 **Van Coppenolle R, Temmerman S. 2019.** A global exploration of tidal wetland creation
 1305 for nature-based flood risk mitigation in coastal cities. *Estuarine, Coastal and Shelf*
 1306 *Science* **226**: 106262
- 1307 **Van Zandt PA, Mopper S. 2002.** Delayed and carryover effects of salinity on flowering in
 1308 *Iris hexagona* (Iridaceae). *American Journal of Botany* **89**: 364–383.
- 1309 **Van Zandt PA, Tobler MA, Mouton E, Hasenstein KH, Mopper S. 2003.** Positive and
 1310 negative consequences of salinity stress for the growth and reproduction of the
 1311 clonal plant, *Iris hexagona*. *Journal of Ecology* **91**: 837–846.
- 1312 **Vasseur DA, DeLong JP, Gilbert B, et al. 2014.** Increased temperature variation poses a
 1313 greater risk to species than climate warming. *Proceedings of the Royal Society B-*
 1314 *Biological Sciences* **281**: 1–8.
- 1315 **Viavattene C, Jiménez JA, Ferreira O, et al. 2018.** Selecting coastal hotspots to storm
 1316 impacts at the regional scale: the Coastal Risk Assessment Framework. *Coastal*
 1317 *Engineering* **134**: 33-47.
- 1318 **Vuik V, Jonkman SN, Borsje BW, Suzuki T. 2016.** Nature-based flood protection: The
 1319 efficiency of vegetated foreshores for reducing wave loads on coastal dikes. *Coastal*
 1320 *Engineering* **116**: 42– 56.
- 1321 **Vuik V, Suh Heo HY, Zhu Z, Borsje BW, Jonkman SN. 2018.** Stem breakage of salt
 1322 marsh vegetation under wave forcing: a field and model study. *Estuarine, Coastal*
 1323 *and Shelf Science* **200**: 41-58.
- 1324 **Waters JM, King TM, Fraser CI, Craw D. 2018.** Crossing the front: contrasting storm-
 1325 forced dispersal dynamics revealed by biological, geological and genetic analysis of
 1326 beach-cast kelp. *Journal of the Royal Society Interface* **15**: 20180046.

- 1327 **White AC, Colmer TD, Cawthray GR, Hanley ME. 2014.** Variable response of three
1328 *Trifolium repens* ecotypes to soil flooding by seawater. *Annals of Botany* **114**: 347-
1329 356.
- 1330 **White E, Kaplan D. 2017.** Restore or retreat? Saltwater intrusion and water management
1331 in coastal wetlands. *Ecosystem Health and Sustainability* **3** e01258.
- 1332 **Williams HFL, Flanagan WM. 2009.** Contribution of Hurricane Rita storm surge
1333 deposition to long-term sedimentation in Louisiana coastal woodlands and marshes.
1334 *Journal of Coastal Research* **56**: 1671-1675.
- 1335 **Zhu Z, Yang Z, Bouma TJ. 2020.** Biomechanical properties of marsh vegetation in space
1336 and time: effects of salinity, inundation and seasonality. *Annals of Botany* 125: doi
1337 10.1093/aob/mcz063.
- 1338 **Zimmerman JK, Hogan JA, Nytch CJ, Bithorn JE. 2018.** Effects of hurricanes and
1339 climate oscillations on annual variation in reproduction in wet forest, Puerto Rico.
1340 *Ecology* **99**: 1402:1410
- 1341 **Zedler JB. 2010.** How frequent storms affect wetland vegetation: a preview of climate-
1342 change impacts. *Frontiers in Ecology and the Environment*, **8**: 540-547.
- 1343

1344 **Figure Legend**

1345 Figure 1. A summary of the principal research priorities (I – IV) and avenues for future
1346 study needed to understand the response of estuarine and coastal plant communities to the
1347 disturbances associated with extreme storm events. The proposed level and overlap of study
1348 (Individual plant, Ecosystem, and Landscape) for each priority is shown. CRAF - Coastal
1349 Flood Risk Frameworks; SDM – Species Distribution Model

1350

1351

Table 1 A summary of the principal acute threats and example responses reported for (semi-)natural coastal plant communities subject to extreme storm events.

Habitat		Threat	Response	Example studies
Sub-tidal	Kelp-forests	Physical damage & dislodgment	Storms cause widespread mortality, but age- and species-specific effects.	Thomsen <i>et al.</i> (2004); Smale and Vance (2016)
	Seagrass	Physical damage	Major losses of seagrass biomass following tropical cyclones.	Sachithanandam <i>et al.</i> (2014); Culliver <i>et al.</i> (2017)
		Sand deposition	High deposition causes (species-specific) mortality.	Cabaco <i>et al.</i> (2008)
		Turbidity	Sediment run-off had greater negative impact than storm damage.	Carlson <i>et al.</i> (2010)
		Rapid salinity change	Long-term, post-storm impacts on community composition.	Ridler <i>et al.</i> (2006); Benjamin <i>et al.</i> (1999)
Inter-tidal	Saltmarsh	Physical damage	Stem breakage likely, although response differs among species. Denudation of vegetation can also occur.	Möller <i>et al.</i> (2014); Vuik <i>et al.</i> (2018); Cahoon (2006)
		Erosion	Storm-induced erosion of the fronting tidal flat may induce marsh erosion and vegetation loss.	Callaghan <i>et al.</i> (2010); Bouma <i>et al.</i> (2016); Leonardi <i>et al.</i> , (2016, 2018)
		Sand, sediment or litter deposition	Burial under sediment or debris can kill vegetation (depending on timing, depth and species).	Callaway and Zedler (2004); Meixler (2017); Leonardi <i>et al.</i> , (2018)
		Changes in salinity or inundation	Heavy rainfall can create opportunities for germination, but salinity changes cause shifts in species and communities.	Zedler (2010); Meixler (2017); Edge <i>et al.</i> , (2020)

	Mangrove	Physical damage/ Erosion	Species-specific variation in tree response (including mortality) to storm damage.	Doyle <i>et al.</i> (1995); Imbert (2018)
			Scour caused <i>Avicennia marina</i> mortality along South African shoreline fringe.	Steinke and Ward (1989)
		Sand/ Litter deposition	Impact of litter largely unknown (see Krauss and Osland 2020), but increased decomposition influences carbon-budgets.	Barr <i>et al.</i> (2012)
			Phosphorus-rich sediments stimulate post-storm forest productivity.	Castañeda-Moya <i>et al.</i> (2010); Adame <i>et al.</i> (2013)
			Sediments covered roots, causing anoxia and tree mortality	Paling <i>et al.</i> (2008)
Supra-tidal	Sand dunes	Physical damage/ Erosion	Sediment loss negatively affects vegetation, but extent depends on dune morphology and vegetation cover.	Hanley <i>et al.</i> (2014); Miller <i>et al.</i> (2015); Schwarz <i>et al.</i> (2019)
		Sand deposition	Sand accumulation induced (species-specific) morphological responses.	Harris <i>et al.</i> (2017); Brown and Zinnert (2018)
		Saline Inundation	Reduced plant performance but species-specific variation in 'stress' responses.	Camprubi <i>et al.</i> , (2012); Hoggart <i>et al.</i> (2014); Hanley <i>et al.</i> (2020a)
	Freshwater marshland	Erosion	Plant mortality facilitated subsequent sediment loss and erosion.	Howes <i>et al.</i> (2010); Hauser <i>et al.</i> (2015)
		Litter deposition	Experimental litter deposition reduced species diversity.	Tate and Battaglia (2013)
		Saline Inundation	Widespread plant mortality observed.	Abbott and Battaglia (2015); Hauser <i>et al.</i> (2015)
	Other habitats	Physical damage	Storm damage caused localised <i>Pinus elliotii</i> mortality in Florida everglades.	Platt <i>et al.</i> (2000)
		Litter deposition	High litter density reduced species	Tate and Battaglia, (2013);

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			diversity in SE USA pine savannah.	Platt <i>et al.</i> (2015)
		Saline Inundation	Negative effects on recovery of Canadian tundra, but with species-specific variation.	Lantz <i>et al.</i> (2015)
			High mortality of Floridian 'freshwater forest' species.	Langston <i>et al.</i> (2017)

1354

I. Reproduction and recruitment

Manipulative experiments to determine impacts of erosion, litter, sedimentation, inundation, flow & waves on;

- Fecundity
- Germination
- Seedling establishment
- Vegetative spread

II. Multiple stressors

Manipulative experiments to determine ecophysiological responses to ACC-linked stressors

- Temperature (averages & extremes)
- Precipitation
- Flooding
- Litter
- Elevated CO₂

III. Community Interactions

Manipulative experiments to elucidate how storms influence post-disturbance;

- Plant competition & facilitation
- Plant-animal interactions
- Plant-microbial interactions
- Soil biogeochemistry

IV. Storm prediction and ecosystem services

Long-term ecological (including remote sensing) monitoring to generate predictive models underpinned by priorities I-III.

- Geomorphological processes
- CRAF
- SDMs

Individual plant

Landscape

Ecosystem